

DESCRIPTION

Title of the Invention

ANTIBODIES FOR DETECTING MICROORGANISMS

5

DESCRIPTION OF BACKGROUND ART

Field of the Invention

The present invention pertains to antibodies useful in the detection of various microorganisms, particularly bacteria, and a method of detecting microorganisms, reagent kits for detection of microorganisms that use said antibodies, and a method for preparing specific antibodies for detecting microorganisms.

Moreover, the present invention is valuable to the drug industry, particularly for diagnostic medicine of microbial infections with an emphasis on bacteria.

Prior Art

Diagnosis of microbial infection is confirmed by detection of the causative pathogen from the infected area or by detection of antibodies to the causative pathogen in serum and body fluids. Detection of the causative pathogen is particularly important in the sense that it makes quick treatment to the patient possible.

Detection of the causative pathogen of infections

can be generally classified as cultivation and identification methods, whereby the causative pathogen is separated and cultivated and then identified based on its biochemical properties;

5 genetic diagnosis, whereby amplification by PCR, etc., is performed based on specific genes of the causative pathogen and thus the causative pathogen is detected; or immunological methods, whereby the causative pathogen is detected using a specific
10 reaction of antibody with surface antigen marker of the causative pathogen. However, it takes time to obtain results by cultivation and identification methods or genetic diagnosis methods. Therefore, diagnosis by immunological methods is commonly used
15 because the causative pathogen can be detected within a short time with high sensitivity and the patient can be quickly and appropriately treated.

Depending on the bacterial species, a variety of marker antigens and antibodies can be used alone or,
20 in combination to detect the causative pathogen of infections by conventional immunological methods.

For instance, it is known that lipopolysaccharide (LPS), which is a genus-specific antigen of *Chlamydia*, is present as an antigen determinant
25 (Stephens, R., et al.: *J. Immunol.*, 128:1083-89, 1982, Caldwell, M.D.: *Inf. Immun.*, 44:306-14, 1984), and antibodies to LPS are used as the reagent

antibody in a variety of diagnostic kits,
particularly for the detection of *Chlamydia*
trachomatis.

Moreover, Ellena M. Peterson et al., (Infection
5 and Immunity, 59(11), 4147-4153, 1991) and Byron E.
Batteiger et al., (Infection and Immunity, 53(3),
530-533, 1986) have reported monoclonal antibodies to
major outer membrane protein (MOMP) of the genus
Chlamydia respectively.

10 Publication of unexamined Japanese patent
application No. 298/1988 discusses an immunodetection
method based on ~~the~~ western blot method that uses a
monoclonal antibody to an approximately 43 kilo
Dalton membrane protein antigen of *Mycoplasma*
15 *pneumoniae*.

Moreover, a method of preparing monoclonal
antibody to *Haemophilus influenzae* and a diagnostic
method that uses said antibody are presented in
Publication of unexamined Japanese patent application
20 No. 148859/1987 (Japanese Patent No. 64065/1994).

Proteins of approximately 20 kilo Dalton isolated
from the sodium cholate extract of the outer membrane
vesicle of *Neisseria gonorrhoeae* strain BS4 (NCTC
11922) are documented and preparation of hybridomas
25 using said substance is disclosed in British Patent
Application No. 2,172,704. Moreover, European Patent
Document no., EP 419238 A1 describes preparation of a

monoclonal antibody, which can bind to a protein of approximately 14 kilo Dalton prepared by using *Neisseria gonorrhoeae* as an immunogen and a method for the preparation of such a monoclonal antibody.

- 5 Additionally, a method of detecting the same *Neisseria gonorrhoeae* using monoclonal antibody to lipopolysaccharide (LPS) is mentioned in Canadian Patent Application No. 1,220,147.

10 However, there are problems with said antibodies and detection methods based on those antibodies in that species specificity to microorganisms is insufficient for proper diagnosis. The antibodies do not detect all serum types as plural surface antigens present within one species.

- 15 Marker antigens used in these prior arts are not standardized so that microorganisms can be detected using a same functional molecule (for instance, protein, LPS or surface polysaccharide component with the same function) which is generally present in
20 cells of various microorganisms and which changes during the course of evolution of the microorganisms as a marker, and an immunodiagnostic method based on the concept of detecting the difference in antigenicity between bacterial species using one
25 molecule as a standard is not known.

DISCLOSURE OF THE INVENTION

The present invention strives to present

antibodies to the same molecule for different microorganisms as a standard marker antigen to make ideal microorganism detection and immunodiagnosis possible, particularly antibodies to the molecular segment in the same intracellular functional component molecule for all microorganisms to be detected that changes during the course of evolution of microorganisms, a method of detecting microorganisms, which is species specific and can cover almost all serum types, a reagent kit for detection of microorganisms that use said antibody, and a method for preparing a specific antibody used for detection of microorganisms.

The present inventors discovered a protein with the same function, preserved in all microorganisms as a useful antigen protein. Usually, it is expected that a structural change of such a protein is extremely small. However, surprisingly it was found that the antigen epitope(s) of this protein has specificity to certain species or genus of microorganisms, and that the antibody for this protein not only has potentialities of being used for specifically identifying species or genus of microorganisms, but also is capable of detecting all serotypes of the target microorganisms.

The inventors focused on intracellular molecule that are present in all microbial cells and differ

between microorganisms in terms of its amino acid sequence, particularly ribosomal protein L7/L12, which is a member of ribosomal proteins. Ribosomal protein L7/L12 is a protein with a molecular weight of approximately 13 kilo Daltons and is known to exist as a ribosomal protein essential in protein synthesis. Progress has been made in understanding the complete amino acid sequence of the protein from several microorganisms including particularly *Escherichia coli* and *Baccillus subtilis*, etc., and 50 % to 65 % homology of the amino acid sequence between the microorganisms has been confirmed.

The inventors focused on the fact that even though there are similarities in said molecule between different microorganisms, this molecule also has a structural segment that is unique to each microorganism and discovered that it is possible to detect various microorganisms with species specificity and to detect all serotypes within the same species by using antibody to the protein. As a result of attempting to develop a technology for immuno-diagnosis of microorganism species using antibody specific to, for instance, *Haemophilus influenzae*, *Streptococcus pneumoniae*, and *Neisseria gonorrhoeae*, the inventors completed the present invention upon discovering that antibody specific to said protein of each species of microorganisms can be

obtained and species-specific detection of different bacteria is possible using said antibody.

The present invention relates to an antibody used for detecting microorganisms, a method of detecting
5 microorganisms using the antibody, a reagent kit for detecting microorganisms using the antibody, and a method for preparing specific antibodies for detecting microorganisms.

1) Antibodies which are antibodies to ribosomal
10 protein of microorganisms and which react specifically with said microorganisms.

2) The antibodies described in 1) above, where the ribosomal protein of the microorganisms is ribosomal protein L7/L12.

15 3) The antibodies described in 1) or 2) above, where said microorganisms are microorganisms, which cause a sexually transmitted disease (STD).

4) The antibodies described in 1) or 2) above, where said microorganisms are microorganisms which
20 cause respiratory tract infection.

5) The antibody described in 4) above, where the causative microorganisms of respiratory tract infection are microorganisms of *Haemophilus influenzae*.

25 6) The antibody described in 4) above, where the causative microorganisms of respiratory tract infection are microorganisms of *Streptococcus*

pneumoniae.

7) The antibody described in 3) above, where the causative microorganisms of sexually transmitted diseases (STD) are microorganisms of *Neisseria gonorrhoeae*.

8) The antibody described in 7) above, which is the antibody to ribosomal protein L7/L12 of *Neisseria gonorrhoeae* and which recognizes a continuous amino acid sequence moiety from 5 to 30 amino acids including the 115th alanine in the amino acid sequence of Sequence ID No. 22 of the Sequence Table.

9) A method of detecting microorganisms, which is characterized by the fact that antibody to intracellular molecules that have the same function for a variety of microorganisms is used.

10) A method of detecting microorganisms, which is characterized by the fact that any antibody described in 1) to 8) above is used.

11) A reagent kit for detecting microorganisms, which is characterized by the fact that antibody to intracellular molecules that have the same function for a variety of microorganisms is used.

12) A reagent kit for detecting microorganisms, which is characterized by the fact that any antibody described in 1) to 8) above is used.

13) A method of preparing any antibody described in 1) to 8) above, characterized by the fact that

ribosomal protein L7/L12 of microorganisms obtained by a gene manipulation procedure or by isolation from microorganisms, peptide moiety thereof, or a synthesized peptide corresponding to the peptide moiety is used as an immunogen.

The present invention will now be explained in detail.

Sub B1
Sequences No. 1 and No. 2 in the Sequence Table are the DNA sequence of the ribosomal protein L7/L12 gene of *Haemophilus influenzae* and corresponding amino acid sequence. Sequences No. 3 and No. 4 are the DNA sequence of the ribosomal protein L7/L12 gene of *Helicobacter pylori* and the corresponding amino acid sequence. Sequences No. 5 and No. 6 show the DNA sequence and the corresponding amino acid sequence of the ribosomal protein L7/L12 gene of *Streptococcus pneumoniae*. Sequences No. 7 and No. 8 show the DNA sequence and the corresponding amino acid sequence of the ribosomal protein L7/L12 gene of *Neisseria gonorrhoeae*. Sequences No. 9 and No. 10 show the DNA sequence and the corresponding amino acid sequence of the ribosomal protein L7/L12 gene of *Neisseria meningitidis*. Sequence No. 11 and Sequence No. 12 in the Sequence Table are the primers DNA for PCR used to acquire the ribosomal protein L7/L12 gene from *Haemophilus influenzae*. Sequences No. 13 and No. 14 in the Sequence Table are the primer DNA for PCR used

to acquire the ribosomal protein L7/L12 gene from *Streptococcus pneumoniae*. Sequences No. 15 and No. 16 in the Sequence Table are the primer DNA for PCR used to acquire the ribosomal protein L7/L12 gene from *Neisseria gonorrhoeae*. Sequences No. 17 and No. 18 show the DNA sequence and the corresponding amino acid sequence of the ribosomal protein L7/L12 gene of *Haemophilus influenzae*. Sequences No. 19 and No. 20 show the DNA sequence and the corresponding amino acid sequence of the ribosomal protein L7/L12 gene of *Streptococcus pneumoniae*. Sequences No. 21 and No. 22 show the DNA sequence and the corresponding amino acid sequence of the ribosomal protein L7/L12 gene of *Neisseria gonorrhoeae*.

Furthermore, the left and right terminals of the amino acid sequences entered in the Sequence Table are amino group (referred to below as the N terminal) and carboxyl group terminals (referred to below as the C terminal), respectively, and the left terminal and right terminal of the base sequence is the 5' terminal and the 3' terminal, respectively.

Moreover, the series of biomolecular experiments of gene preparation mentioned in this text can be performed by methods entered in standard experimental manuals. "Molecular cloning: A laboratory manual," Cold Spring Harbor Laboratory Press, Sambrook, J. et al. (1989), is an example of the aforementioned

standard experimental manual.

The term microorganism in the present invention refers to all species of microorganisms, including bacteria, yeast, mold, *Actinomyces*, *rickettsia*, etc., but bacteria in particular, pose a problem in terms of diagnosis of microbial infections.

The term "antibody which reacts specifically with microorganisms" as used in the present invention means an antibody, which reacts specifically with a species or group of microorganisms. An antibody, which reacts specifically with a species of microorganisms, is particularly useful for the diagnosis of microbial infection diseases.

In the present invention, causative microorganisms of STD (sexually transmitted disease) include, but are not limited to, *Neisseria gonorrhoeae*, *Chlamydia trachomatis*, *Candida albicans*, *Treponema pallidum*, and *Ureaplasma urealyticum*.

In the present invention, causative microorganisms of respiratory tract infection include, but are not limited to, *Haemophilus influenzae*, *Streptococcus pneumoniae*, *Chlamydia pneumoniae*, *Mycoplasma pneumoniae*, *Klebsiella pneumoniae*, *Staphylococcus aureus*, *Pseudomonas aeruginosa*, *Streptococcus sp. Group-A*, *Mycobacterium tuberculosis*, *Legionella pneumophila*, and *Aspergillus spp.*

The term antibody in the present invention means a polyclonal antibody or monoclonal antibody that can be made using the entire length or only a partial peptide of said ribosomal protein. Although there are
5 no special restrictions to the peptide length for making the antibody, the segment should be of the length characterizing the ribosomal protein L7/L12, and a peptide of 5 amino acids or longer, particularly 8 amino acids or longer, is preferred.

10 Antiserum containing antibody (polyclonal antibody) that identifies ribosomal protein L7/L12 can be obtained by inoculating laboratory animals with adjuvant and a peptide or the full length protein as it is or when necessary, after being cross linked
15 with a carrier protein such as KLH (keyhole-limpet hemocyanin) and BSA (bovine serum albumin) and recovering the serum. Moreover, the antibody can be used after it has been purified from the antiserum. The laboratory animals that are inoculated include
20 sheep, horses, goats, rabbits, mice, rats, etc., and sheep, rabbits, etc., are particularly preferred for preparation of polyclonal antibody. Moreover, monoclonal antibody can also be obtained by
25 mice are preferred in this case. The entire length of the said protein, or its partial peptide consisting amino acid residues of 5 or more,

preferably 8 or more, residues that has been fused with GST (glutathione S-transferase), etc., can be purified and used as antigen, or it can be used as antigen without being purified. The antibody can
5 also be the genetic recombination antibody expressed cellularly using immunoglobulin genes that have been separated by a variety of methods in manuals ("Antibodies: A Laboratory manual," E. Harlow et al., Cold Spring Harbor Laboratory), cloning methods,
10 etc.

Antibody to ribosomal protein L7/L12 that can be employed as the marker antigen of the present invention can be obtained by the following 3 methods, and other similar methods as well. The methods,
15 however, are not limited to these.

a) The desired antibody can be acquired by synthesizing a peptide fragment consisting of 5-30 amino acids for microorganisms with a known ribosomal protein L7/L12 genetic sequence and amino acid
20 sequence using the region least similar to the amino acid sequence of said protein of another microorganisms and making polyclonal antibody, or monoclonal antibody, using this peptide fragment as the immunogen.

25 Moreover, it is possible to acquire the entire sequence of said gene by using a conventional genetic procedure, such as gene amplification by PCR with the

DNA sequence at both terminals of said known genetic sequence as the probe, or hybridization using the sequence of a homologous segment as the template probe.

5 Then, a fused gene with another protein gene is constructed and said fused gene is inserted into the host by conventional gene insertion methods using *Escherichia coli*, etc., as the host and expressed in large quantities. The desired protein antigen can
10 then be acquired by purifying the expressed protein by affinity column methods with antibody to the protein that was used as the fusion protein. In such a case, even if antibody to the amino acid segment conserved within microorganisms is acquired, it does
15 not coincide with the purpose of the present invention because the full length of ribosomal protein L7/L12 becomes the antigen. Consequently, hybridoma that produces monoclonal antibody to the antigen that has been obtained by this method is
20 acquired by conventional methods and the desired antibody can be obtained by selecting a clone that produces antibody that will react only with the desired microorganisms.

b) First, since there is 50 to 60% homology
25 between bacterial species in terms of their ribosomal protein L7/L12 amino acid sequence, it is possible to easily acquire said protein genes for microorganisms

with unknown L7/L12 amino acid sequence by conventional genetic procedures, such as gene amplification of a specific sequence segment by PCR methods based on the sequence of the homologous segments of the amino acid sequence, or hybridization with the homologous segments as the template probe, using bacteria having a known ribosomal protein L7/L12 amino acid sequence.

Then, a fused gene with another protein gene is constructed and said fused gene is inserted into the host by conventional gene insertion methods using *Escherichia coli*, etc., as the host and expressed in large quantities. The desired protein antigen can then be acquired by purifying the expressed protein by affinity column methods with antibody to the protein that was used as the fusion protein. In such a case, even if antibody to the amino acid segment conserved within microorganisms is acquired, it does not coincide with the purpose of the present invention because the full length of ribosomal protein L7/L12 becomes the antigen. Consequently, hybridoma that produces monoclonal antibody to the antigen that has been obtained by this method is acquired by conventional methods and the desired antibody can be obtained by selecting a clone that produces antibody that will react only with the desired microorganisms.

c) Ribosomal protein L7/L12 that has been purified to a high purity can also be obtained by another method, in the case where the amino acid sequence of the ribosomal protein L7/L12 is unknown
5 whereby a peptide of 5 to 30 amino acids corresponding to the common sequence segment retained in the microorganisms is synthesized from the known amino acid sequence of the ribosomal protein L7/L12, and polyclonal antibody or monoclonal antibody to
10 this peptide sequence is made by conventional methods. Then the highly purified Ribosomal protein L7/L12 from disrupted microorganisms is obtained by affinity column chromatography using said antibody.

If purity of the protein is insufficient, it can
15 be purified by conventional methods, such as ion exchange chromatography, hydrophobic chromatography, gel filtration, etc., after which the eluted fraction of ribosomal protein L7/L12 is identified by western blotting, etc., using antibody that was made to
20 obtain the pure fraction. The desired antibody can be obtained by acquiring hybridoma or polyclonal antibody by conventional methods using the pure ribosomal protein L7/L12 antigen that has been obtained and selecting hybridoma or polyclonal
25 antibody that will react specifically with the desired bacteria, as in b).

The antibody of the present invention specific to

a variety of microorganisms that has been obtained by the methods in a) through c), etc., can be used in a variety of immunoassay methods to obtain diagnostic reagent kits specific to a variety of microorganisms.

5 For example, this antibody can be used in aggregation reactions where antibody is adsorbed on polystyrene latex particles, ELISA, which is a conventional technology performed in a microtiter plate, conventional immunochromatography methods, sandwich
10 assay, whereby said antibody labeled with colored particles or particles that have coloring capability, or with enzyme or phosphor, and magnetic microparticles coated with capture antibody, etc., are used, etc.

15 The term detection methods for microorganism using antibody means detection methods that use conventional immunoassay such as aggregation reactions where antibody is adsorbed on polystyrene latex particles, ELISA, which is a conventional
20 technology performed in a microtiter plate, conventional immunochromatography methods, sandwich assay, whereby said antibody labeled with colored particles or particles that have coloring capability, or with enzyme or phosphor, and magnetic
25 microparticles coated with capture antibody, etc., are used, etc.

Moreover, the optical immunoassay (OIA)

technology described in International Patent Application Japanese Laid-open (Toku-Hyou) No. 509565/1995, in which microorganisms are detected by an optical interference induced by an antibody
5 reaction on the optical thin film which is formed by silicone or silicon nitride, is a useful detection method using an antibody.

Moreover, treatment with an extraction reagent that uses a variety of surfactants, beginning with
10 Triton X-100 and Tween-20, enzyme treatment with an appropriate enzyme, such as protease, etc., known methods whereby the cell structure is disrupted, beginning with disruption of the microorganism by physical methods, can be used to extract the
15 intracellular marker antigen from the necessary microorganism in said detection method. However, it is preferred that the extraction conditions be designed using a combination of surfactants, etc., so that the conditions are optimized for extraction of
20 each microorganism with reagents.

Moreover, the term a reagent kit for detection of microorganisms using antibody means a reagent kit that uses said detection method.

For instance, in the case of obtaining the
25 specific antibody to *Haemophilus influenzae*, which is of extreme diagnostic significance as a causative pathogen of pneumonia, bronchitis, meningitis, etc.,

the amino acid sequence and DNA sequence of ribosomal protein L7/L12 is entered in data bases, etc.

1 The amino acid and DNA sequence of ribosomal protein L7/L12 of *Haemophilus influenzae* are shown in
5 "Sequences No. 1 and No. 2."

1 Consequently, in the case of this bacteria, it is possible to similarly compare the amino acid sequence of ribosomal protein L7/L12 with the same protein of, for instance, *Helicobacter pylori*, which is shown in
10 "Sequence No. 3 and No. 4," and synthesize a peptide of 5 to 30 amino acids for the segment of low homology and make polyclonal antibody or monoclonal antibody specific to *Haemophilus influenzae* using this peptide.

15 In the case of a specific polyclonal antibody, it is preferred that IgG fraction be obtained by purification of the antiserum of immunized laboratory animals with a protein A column, etc., and affinity purification be performed with the synthetic peptide used in immunization of the laboratory animals.

20 Moreover, PCR primers based on the sequences of N-terminal and C-terminal, for example, the PCR primers shown in Sequences No. 11 and No. 12 in the Sequence Table, were designed from the DNA sequence
25 of ribosomal protein L7/L12 of *Haemophilus influenzae*. Utilizing homology of the PCR primers, DNA fragments amplified by the PCR method or the like

using genomic DNA which is extracted from cultivated *Haemophilus influenzae* can be acquired by a conventional method. The entire length of the gene for ribosomal protein L7/L12 of *Haemophilus influenzae* can be acquired by the analysis of the DNA sequence information of these fragments.

The ribosomal protein L7/L12 gene of *Haemophilus influenzae* thus acquired forms a fusion protein gene with, for instance, GST etc., and expression vector is constructed using the appropriate expression plasmid, *Escherichia coli* is transformed and large quantities of said protein can be expressed. A suitable amount of the transformed *Escherichia coli* is cultivated and the crushed bacterial fluid that is recovered is purified by an affinity column using GST to obtain the ribosomal protein L7/L12 and GST fusion protein of *Haemophilus influenzae*. It is also possible to acquire the target specific monoclonal antibody by establishing plural hybridomas using said protein as it is or GST moiety fragments cut from the protein by protease or the like, as an antigen protein and selecting the antibody which exhibits a specific response to *Haemophilus influenzae* bacteria, or a disrupt fluid of the bacteria, or ribosomal protein L7/L12 of *Haemophilus influenzae*.

Moreover, the amino acid sequence and the DNA sequence of ribosomal protein L7/L12 of *Streptococcus*

pneumoniae which is also highly significant as a diagnostic agent for respiratory infection diseases as well as *Haemophilus influenzae*, are known from descriptions in data bases and the like. The amino acid and DNA sequences of ribosomal protein L7/L12 of *Streptococcus pneumoniae* are shown in Sequences No. 5 and No. 6 of sequence table.

1WS
36
It is therefore possible to acquire a polyclonal antibody or monoclonal antibody which is specific to *Streptococcus pneumoniae* by designing a PCR primer, the PCR primer shown by Sequence ID No. 13 or 14 in the Sequence Table, for example, based on the sequences of N-terminal and C-terminal of DNA sequence of Ribosomal Proteins L7/L12 of *Streptococcus pneumoniae* in the same manner as in the case of *Haemophilus influenzae*, and processing thereafter in the same manner as in the case of *Haemophilus influenzae*.

The hybridoma AMSP-2 which produces the monoclonal antibody specific to *Streptococcus pneumoniae* has been deposited with the National Institute of Bioscience and Human-Technology, the Agency of Industrial Science and Technology, the Ministry of International Trade and Industry, Japan, on July 28, 1999, with the deposition number FERM BP-6807.

Moreover, although the DNA and amino acid

sequences of the ribosomal protein L7/L12 of, for instance, *Neisseria gonorrhoeae*, which is the causative pathogen of gonorrhea and has been shown to have diagnostic significance as a typical causative pathogen of STD, were unknown, a major part of the DNA sequence and amino acid sequence, which was determined by the *Neisseria Gonorrhea Genome Project* at Oklahoma University, USA, is disclosed on the Internet.

When part of the known DNA sequence of ribosomal protein L7/L12 was used to probe the existence of DNA fragments with a similar sequence, it was found that DNA sequence corresponding to the ribosomal protein L7/L12 gene is present and it was possible to obtain data on its entire DNA sequence. The entire base sequence and corresponding amino acid sequence of the ribosomal protein L7/L12 gene of this *Neisseria gonorrhoeae* are shown in Sequences No. 7 and No. 8 of the sequence table.

It is therefore possible to acquire the target antibody which is specific to *Neisseria gonorrhoeae* having the entire or partial Ribosomal Protein L7/L12 of *Neisseria gonorrhoeae* as an antigen by designing a PCR primer, the PCR primer shown by Sequence ID No. 15 or 16 in the Sequence Table, for example, based on the sequences of N-terminal and C-terminal of DNA sequence of Ribosomal Protein L7/L12 of *Neisseria*

gonorrhoeae in the same manner as in the case of
Haemophilus influenzae, and *Streptococcus pneumoniae*,
and processing thereafter in exactly the same manner
as in the case of *Haemophilus influenzae* or
5 *Streptococcus pneumoniae*.

LWS
BG
1
2
3
4
5
6
7
8
9
10
11
12
13
14
15
16
17
18
19
20
21
22
23
24
25
26
27
28
29
30
31
32
33
34
35
36
37
38
39
40
41
42
43
44
45
46
47
48
49
50
51
52
53
54
55
56
57
58
59
60
61
62
63
64
65
66
67
68
69
70
71
72
73
74
75
76
77
78
79
80
81
82
83
84
85
86
87
88
89
90
91
92
93
94
95
96
97
98
99
100
101
102
103
104
105
106
107
108
109
110
111
112
113
114
115
116
117
118
119
120
121
122
123
124
125
126
127
128
129
130
131
132
133
134
135
136
137
138
139
140
141
142
143
144
145
146
147
148
149
150
151
152
153
154
155
156
157
158
159
160
161
162
163
164
165
166
167
168
169
170
171
172
173
174
175
176
177
178
179
180
181
182
183
184
185
186
187
188
189
190
191
192
193
194
195
196
197
198
199
200
201
202
203
204
205
206
207
208
209
210
211
212
213
214
215
216
217
218
219
220
221
222
223
224
225
226
227
228
229
230
231
232
233
234
235
236
237
238
239
240
241
242
243
244
245
246
247
248
249
250
251
252
253
254
255
256
257
258
259
260
261
262
263
264
265
266
267
268
269
270
271
272
273
274
275
276
277
278
279
280
281
282
283
284
285
286
287
288
289
290
291
292
293
294
295
296
297
298
299
300
301
302
303
304
305
306
307
308
309
310
311
312
313
314
315
316
317
318
319
320
321
322
323
324
325
326
327
328
329
330
331
332
333
334
335
336
337
338
339
340
341
342
343
344
345
346
347
348
349
350
351
352
353
354
355
356
357
358
359
360
361
362
363
364
365
366
367
368
369
370
371
372
373
374
375
376
377
378
379
380
381
382
383
384
385
386
387
388
389
390
391
392
393
394
395
396
397
398
399
400
401
402
403
404
405
406
407
408
409
410
411
412
413
414
415
416
417
418
419
420
421
422
423
424
425
426
427
428
429
430
431
432
433
434
435
436
437
438
439
440
441
442
443
444
445
446
447
448
449
450
451
452
453
454
455
456
457
458
459
460
461
462
463
464
465
466
467
468
469
470
471
472
473
474
475
476
477
478
479
480
481
482
483
484
485
486
487
488
489
490
491
492
493
494
495
496
497
498
499
500
501
502
503
504
505
506
507
508
509
510
511
512
513
514
515
516
517
518
519
520
521
522
523
524
525
526
527
528
529
530
531
532
533
534
535
536
537
538
539
540
541
542
543
544
545
546
547
548
549
550
551
552
553
554
555
556
557
558
559
560
561
562
563
564
565
566
567
568
569
570
571
572
573
574
575
576
577
578
579
580
581
582
583
584
585
586
587
588
589
590
591
592
593
594
595
596
597
598
599
600
601
602
603
604
605
606
607
608
609
610
611
612
613
614
615
616
617
618
619
620
621
622
623
624
625
626
627
628
629
630
631
632
633
634
635
636
637
638
639
640
641
642
643
644
645
646
647
648
649
650
651
652
653
654
655
656
657
658
659
660
661
662
663
664
665
666
667
668
669
670
671
672
673
674
675
676
677
678
679
680
681
682
683
684
685
686
687
688
689
690
691
692
693
694
695
696
697
698
699
700
701
702
703
704
705
706
707
708
709
710
711
712
713
714
715
716
717
718
719
720
721
722
723
724
725
726
727
728
729
730
731
732
733
734
735
736
737
738
739
740
741
742
743
744
745
746
747
748
749
750
751
752
753
754
755
756
757
758
759
760
761
762
763
764
765
766
767
768
769
770
771
772
773
774
775
776
777
778
779
780
781
782
783
784
785
786
787
788
789
790
791
792
793
794
795
796
797
798
799
800
801
802
803
804
805
806
807
808
809
810
811
812
813
814
815
816
817
818
819
820
821
822
823
824
825
826
827
828
829
830
831
832
833
834
835
836
837
838
839
840
841
842
843
844
845
846
847
848
849
850
851
852
853
854
855
856
857
858
859
860
861
862
863
864
865
866
867
868
869
870
871
872
873
874
875
876
877
878
879
880
881
882
883
884
885
886
887
888
889
890
891
892
893
894
895
896
897
898
899
900
901
902
903
904
905
906
907
908
909
910
911
912
913
914
915
916
917
918
919
920
921
922
923
924
925
926
927
928
929
930
931
932
933
934
935
936
937
938
939
940
941
942
943
944
945
946
947
948
949
950
951
952
953
954
955
956
957
958
959
960
961
962
963
964
965
966
967
968
969
970
971
972
973
974
975
976
977
978
979
980
981
982
983
984
985
986
987
988
989
990
991
992
993
994
995
996
997
998
999
1000
1001
1002
1003
1004
1005
1006
1007
1008
1009
1010
1011
1012
1013
1014
1015
1016
1017
1018
1019
1020
1021
1022
1023
1024
1025
1026
1027
1028
1029
1030
1031
1032
1033
1034
1035
1036
1037
1038
1039
1040
1041
1042
1043
1044
1045
1046
1047
1048
1049
1050
1051
1052
1053
1054
1055
1056
1057
1058
1059
1060
1061
1062
1063
1064
1065
1066
1067
1068
1069
1070
1071
1072
1073
1074
1075
1076
1077
1078
1079
1080
1081
1082
1083
1084
1085
1086
1087
1088
1089
1090
1091
1092
1093
1094
1095
1096
1097
1098
1099
1100
1101
1102
1103
1104
1105
1106
1107
1108
1109
1110
1111
1112
1113
1114
1115
1116
1117
1118
1119
1120
1121
1122
1123
1124
1125
1126
1127
1128
1129
1130
1131
1132
1133
1134
1135
1136
1137
1138
1139
1140
1141
1142
1143
1144
1145
1146
1147
1148
1149
1150
1151
1152
1153
1154
1155
1156
1157
1158
1159
1160
1161
1162
1163
1164
1165
1166
1167
1168
1169
1170
1171
1172
1173
1174
1175
1176
1177
1178
1179
1180
1181
1182
1183
1184
1185
1186
1187
1188
1189
1190
1191
1192
1193
1194
1195
1196
1197
1198
1199
1200
1201
1202
1203
1204
1205
1206
1207
1208
1209
1210
1211
1212
1213
1214
1215
1216
1217
1218
1219
1220
1221
1222
1223
1224
1225
1226
1227
1228
1229
1230
1231
1232
1233
1234
1235
1236
1237
1238
1239
1240
1241
1242
1243
1244
1245
1246
1247
1248
1249
1250
1251
1252
1253
1254
1255
1256
1257
1258
1259
1260
1261
1262
1263
1264
1265
1266
1267
1268
1269
1270
1271
1272
1273
1274
1275
1276
1277
1278
1279
1280
1281
1282
1283
1284
1285
1286
1287
1288
1289
1290
1291
1292
1293
1294
1295
1296
1297
1298
1299
1300
1301
1302
1303
1304
1305
1306
1307
1308
1309
1310
1311
1312
1313
1314
1315
1316
1317
1318
1319
1320
1321
1322
1323
1324
1325
1326
1327
1328
1329
1330
1331
1332
1333
1334
1335
1336
1337
1338
1339
1340
1341
1342
1343
1344
1345
1346
1347
1348
1349
1350
1351
1352
1353
1354
1355
1356
1357
1358
1359
1360
1361
1362
1363
1364
1365
1366
1367
1368
1369
1370
1371
1372
1373
1374
1375
1376
1377
1378
1379
1380
1381
1382
1383
1384
1385
1386
1387
1388
1389
1390
1391
1392
1393
1394
1395
1396
1397
1398
1399
1400
1401
1402
1403
1404
1405
1406
1407
1408
1409
1410
1411
1412
1413
1414
1415
1416
1417
1418
1419
1420
1421
1422
1423
1424
1425
1426
1427
1428
1429
1430
1431
1432
1433
1434
1435
1436
1437
1438
1439
1440
1441
1442
1443
1444
1445
1446
1447
1448
1449
1450
1451
1452
1453
1454
1455
1456
1457
1458
1459
1460
1461
1462
1463
1464
1465
1466
1467
1468
1469
1470
1471
1472
1473
1474
1475
1476
1477
1478
1479
1480
1481
1482
1483
1484
1485
1486
1487
1488
1489
1490
1491
1492
1493
1494
1495
1496
1497
1498
1499
1500
1501
1502
1503
1504
1505
1506
1507
1508
1509
1510
1511
1512
1513
1514
1515
1516
1517
1518
1519
1520
1521
1522
1523
1524
1525
1526
1527
1528
1529
1530
1531
1532
1533
1534
1535
1536
1537
1538
1539
1540
1541
1542
1543
1544
1545
1546
1547
1548
1549
1550
1551
1552
1553
1554
1555
1556
1557
1558
1559
1560
1561
1562
1563
1564
1565
1566
1567
1568
1569
1570
1571
1572
1573
1574
1575
1576
1577
1578
1579
1580
1581
1582
1583
1584
1585
1586
1587
1588
1589
1590
1591
1592
1593
1594
1595
1596
1597
1598
1599
1600
1601
1602
1603
1604
1605
1606
1607
1608
1609
1610
1611
1612
1613
1614
1615
1616
1617
1618
1619
1620
1621
1622
1623
1624
1625
1626
1627
1628
1629
1630
1631
1632
1633
1634
1635
1636
1637
1638
1639
1640
1641
1642
1643
1644
1645
1646
1647
1648
1649
1650
1651
1652
1653
1654
1655
1656
1657
1658
1659
1660
1661
1662
1663
1664
1665
1666
1667
1668
1669
1670
1671
1672
1673
1674
1675
1676
1677
1678
1679
1680
1681
1682
1683
1684
1685
1686
1687
1688
1689
1690
1691
1692
1693
1694
1695
1696
1697
1698
1699
1700
1701
1702
1703
1704
1705
1706
1707
1708
1709
1710
1711
1712
1713
1714
1715
1716
1717
1718
1719
1720
1721
1722
1723
1724
1725
1726
1727
1728
1729
1730
1731
1732
1733
1734
1735
1736
1737
1738
1739
1740
1741
1742
1743
1744
1745
1746
1747
1748
1749
1750
1751
1752
1753
1754
1755
1756
1757
1758
1759
1760
1761
1762
1763
1764
1765
1766
1767
1768
1769
1770
1771
1772
1773
1774
1775
1776
1777
1778
1779
1780
1781
1782
1783
1784
1785
1786
1787
1788
1789
1790
1791
1792
1793
1794
1795
1796
1797
1798
1799
1800
1801
1802
1803
1804
1805
1806
1807
1808
1809
1810
1811
1812
1813
1814
1815
1816
1817
1818
1819
1820
1821
1822
1823
1824
1825
1826
1827
1828
1829
1830
1831
1832
1833
1834
1835
1836
1837
1838
1839
1840
1841
1842
1843
1844
1845
1846
1847
1848
1849
1850
1851
1852
1853
1854
1855
1856
1857
1858
1859
1860
1861
1862
1863
1864
1865
1866
1867
1868
1869
1870
1871
1872
1873
1874
1875
1876
1877
1878
1879
1880
1881
1882
1883
1884
1885
1886
1887
1888
1889
1890
1891
1892
1893
1894
1895
1896
1897
1898
1899
1900
1901
1902
1903
1904
1905
1906
1907
1908
1909
1910
1911
1912
1913
1914
1915
1916
1917
1918
1919
1920
1921
1922
1923
1924
1925
1926
1927
1928
1929
1930
1931
1932
1933
1934
1935
1936
1937
1938
1939
1940
1941
1942
1943
1944
1945
1946
1947
1948
1949
1950
1951
1952
1953
1954
1955
1956
1957
1958
1959
1960
1961
1962
1963
1964
1965
1966
1967
1968
1969
1970
1971
1972
1973
1974
1975
1976
1977
1978
1979
1980
1981
1982
1983
1984
1985
1986
1987
1988
1989
1990
1991
1992
1993
1994
1995
1996
1997
1998
1999
2000
2001
2002
2003
2004
2005
2006
2007
2008
2009
2010
2011
2012
2013
2014
2015
2016
2017
2018
2019
2020
2021
2022
2023
2024
2025
2026
2027
2028
2029
2030
2031
2032
2033
2034
2035
2036
2037
2038
2039
2040
2041
2042
2043
2044
2045
2046
2047
2048
2049
2050
2051
2052
2053
2054
2055
2056
2057
2058
2059
2060
2061
2062
2063
2064
2065
2066
2067
2068
2069
2070
2071
2072
2073
2074
2075
2076
2077
2078
2079
2080
2081
2082
2083
2084
2085
2086
2087
2088
2089
2090
2091
2092
2093
2094
2095
2096
2097
2098
2099
2100
2101
2102
2103
2104
2105
2106
2107
2108
2109
2110
2111
2112
2113
2114
2115
2116
2117
2118
2119
2120
2121
2122
2123
2124
2125
2126
2127
2128
2129
2130
2131
2132
2133
2134
2135
2136
2137
2138
2139
2140
2141
2142
2143
2144
2145
2146
2147
2148
2149
2150
2151
2152
2153
2154
2155
2156
2157
2158
2159
2160
2161
2162
2163
2164
2165
2166
2167
2168
2169
2170
2171
2172
2173
2174
2175
2176
2177
2178
2179
2180
2181
2182
2183
2184
2185
2186
2187
2188
2189
2190
2191
2192
2193
2194
2195
2196
2197
2198
2199
2200
2201
2202
2203
2204
2205
2206
2207
2208
2209
2210
2211
2212
2213
2214
2215
2216
2217
2218
2219
2220
2221
2222
2223
22

be used in all known types of immunoassay, such as aggregation whereby said antibody is adsorbed on polystyrene latex, ELISA, which is a conventional technology performed in a microtiter plate, conventional immunochromatography, sandwich assay, whereby said antibody labeled with colored particles or particles that have coloring capability, or enzymes or phosphor, and magnetic particles coated with capture antibody are used, etc.

10 Moreover, antibody that is made based on the present invention can simultaneously function in any of these immunoassay methods as a so-called capture antibody that captures said antigen protein in solid or liquid phase and a so-called enzyme-labeled antibody by modification using an enzyme, such as 15 peroxidase and alkali phosphatase, etc., by conventional methods.

BEST MODE FOR CARRYING OUT THE INVENTION

20 The following examples are given to explain the present invention in actual terms, the present invention not being restricted to these examples.

Example 1

Cloning of ribosomal protein L7/L12 genes from
Haemophilus influenzae

1 After inoculating an appropriate amount of *Haemophilus influenzae* strain ATCC9334 (IID984) (obtained from Tokyo University School of Medicine

Laboratories) in a chocolate agar culture medium, the strain was cultivated for 24 hours in a CO₂ incubator under conditions of 37 °C and 0.5 <5.0> % CO₂. The colonies that grew were suspended in a TE buffer (manufactured by Wako Pure Chemical Co., Ltd.) to a final concentration of approximately 5 x 10⁹ CFU/ml. Approximately 1.5 ml of this suspension was transferred to a microcentrifuge tube and centrifuged for 2 minutes at 10,000 rpm. The supernatant was discarded. The sediment was resuspended in 567 µl TE buffer. Then 30 µl 10 % sodium dodecylsulfate (SDS) and 3 µl 20 mg/ml Proteinase K solution were added and thoroughly mixed. The suspension was incubated for another hour at 37 °C. Next, after adding 80 µl 10 % cetyl trimethyl ammonium bromide/0.7 M NaCl solution and thoroughly mixing the product, it was incubated for 10 minutes at 65 °C. Next, 700 µl chloroform-isoamyl alcohol solution at a volume ratio of 24:1 was added and stirred well. The solution was centrifuged for 5 minutes (while being kept at 4 °C) at 12,000 rpm using a microcentrifugation device and the aqueous fraction was transferred to a new microtube. Isopropanol was added to the fraction at 0.6-times its volume and the tube was vigorously shaken to form sediment of the DNA. The white DNA sediment was scooped with a glass rod and transferred to a different microcentrifuge tube containing 1 ml

70 % ethanol (cooled to -20 °C).

Next, the product was centrifuged for 5 minutes at 10,000 rpm and the supernatant was gently removed. Then another 1 ml 70 % ethanol was added and the product was centrifuged for 5 more minutes.

Once the supernatant had been removed, the sediment was dissolved in 100 µl TE buffer to obtain the DNA solution. The concentration of the genomic DNA solution was determined quantitatively according to E5, Spectrophotometric determination of the amount of DNA or RNA "Molecular cloning: A laboratory manual," 1989, Eds. Sambrook, J., Fritsch, E.F., and Maniatis, T., Cold Spring Harbor Laboratory Press.

PCR (polymerase chain reaction) was performed using 10 ng of this genomic DNA. Taq polymerase (Takara Co., ltd., code R001A) was employed for PCR. Then 5 µl of buffer, 4 µl dNTP mixture, and 200 pmol each of synthetic oligonucleotide-A, shown in Sequence No. 11 of the Sequence Table and synthetic oligonucleotide-B, shown in Sequence No. 12 of the Sequence table were added to the enzyme to bring the final volume to 50 µl.

This mixture was cycled 5 times with a TaKaRa PCR Thermal Cycler 480 for 1 minute at 95 °C, 2 minutes at 50 °C, and 3 minutes at 72 °C and was then cycled 25 times for 1 minute at 95 °C, 2 minutes at 60 °C, and 3 minutes at 72 °C. Electrophoresis was performed

in 1.5 % agarose gel using some of this PCR product. This product was then stained with ethidium bromide (Nihon Gene Co., ltd.) and observed under ultraviolet rays to confirm amplification of approximately 400 bp
5 cDNA. After digestion treatment with restriction endonucleases BamHI and XhoI, electrophoresis was performed in 1.5 % agarose gel and staining with ethidium bromide was carried out. An approximately 370 bp band was cut out from the gel. This band was
10 purified with Suprec01 (Takara Co., Ltd.) and then inserted into pGEX-4T-1 (Pharmacia), which is a commercial vector. This same vector can function as an expression vector for the desired molecule, which can express fused protein with GST protein, by
15 insertion of the desired gene fragment into the appropriate restriction endonuclease site.

Actually, vector pGEX-4T-1 and the previous DNA were mixed together at a molar ratio of 1:3 and DNA was inserted into the vector with T4 DNA ligase
20 (Invitrogen Co.). Vector pGEX-4T-1 into which DNA had been inserted was genetically introduced to *Escherichia coli* one-shot competent cells (Invitrogen Co., Ltd.) and then inoculated in a plate of L-Broth (Takara Co., Ltd.) semi-solid culture plate containing
25 50 µg/ml ampicillin (Sigma). The plate was then set-aside at 37 °C for 12 hours and the colonies that grew were selected at random and inoculated into 2 ml

L-Broth liquid culture medium containing the same concentration of ampicillin. Shake cultivation was performed at 37 °C for 8 hours and the bacteria were recovered and the plasmid was separated using Wizard
5 Miniprep (Promega) in accordance with the attached literature. The plasmid was cleaved with restriction endonuclease BamHI/XhoI. Insertion of said PCR product was confirmed by cutting out approximately 370 bp DNA. The base sequence of the DNA that had
10 been inserted was determined using said clone.

125
B11 → (Determination of the base sequence of the inserted DNA fragment was performed using the Fluorescence Sequencer of Applied Biosystems. The sequence sample was prepared using PRISM, Ready
15 Reaction Dye Terminator Cycle Sequencing Kit (Applied Biosystems). First, 9.5 µl reaction stock solution, 4.0 µl T7 promoter primer at 0.8 pmol/µl (Gibco BRL) and 6.5 µl of template DNA for sequencing at 0.16 µg/µl were added to a microtube with a capacity of
20 0.5 ml, mixed and superposed with 100 µl mineral oil. PCR amplification was performed for 25 cycles, where one cycle consisted of 30 seconds at 96 °C, 15 seconds at 55 °C, and 4 minutes at 60 °C. The product was then kept at 4 °C for 5 minutes. After the
25 reaction was completed, 80 µl sterilized pure water was added and stirred. The product was centrifuged and the aqueous layer was extracted 3 times with

phenol-chloroform. Ten ml <microliters> 3 M sodium acetate (pH 5.2) and 300 µl ethanol were added to 100 µl aqueous layer and stirred. The product was then centrifuged for 15 minutes at room temperature and 14,000 rpm and the sediment was recovered. Once the sediment was washed with 75 % ethanol, it was dried under a vacuum for 2 minutes to obtain the sequencing sample. The sequencing sample was dissolved in formamide containing 4 µl 10 mM EDTA and denatured for 2 minutes at 90 °C. This was then cooled in ice and applied to sequencing.

One of the 5 clones obtained had homology of the sequence with the probe used for PCR. In addition, DNA sequences extremely similar to the gene sequence of ribosomal protein L7/L12 gene of the other microorganisms, for example, *Neisseria gonorrhoeae*, were discovered. The entire base sequence and the corresponding amino acid sequence of the structural gene moiety are as shown in Sequence ID No. 17 and No. 18 of the Sequence Table. This gene fragment clearly codes for *Haemophilus influenzae* ribosomal protein L7/L12.

Example 2

Mass expression in *Escherichia coli* and purification of ribosomal protein L7/L12 from *Haemophilus influenzae*

Fifty milliliters *Escherichia coli* into which

expression vector had been inserted were cultivated overnight in LB at 37 °C. Then 500 ml YT medium at concentration that was twice that of the
aforementioned culture was heated at 37 °C for 1
5 hour. Fifty milliliters of the *Escherichia coli* solution that had been cultivated overnight were introduced to 500 ml of the aforementioned medium. One hour later, 550 µl 100 mM isopropyl β-(D)-thiogalactopyranoside (IPTG) were introduced and
10 cultivated for 4 hours. The product was then recovered and introduced to centrifugation tube at each 250 ml and centrifuged for 10 minutes at 7,000 rpm.

The supernatant was discarded and dissolved in 25
15 ml each 50 mM Tris-HCl at a pH of 7.4 and Lysis buffer containing 25 % sucrose.

Furthermore, 1.25 ml 10 % Nonidet P-40 (NP-40) and 125 µl 1 M MgCl₂ were added and transferred to a plastic tube. Sonication was performed 1 minute x 5
20 times while ice cold. The product was centrifuged for 15 minutes at 12,000 rpm and the supernatant was recovered.

Next, the aforementioned supernatant was adsorbed on a glutathione agarose column conditioned with
25 phosphate-buffered saline (PBS).

Then the column was washed with twice the bed volume using a washing solution containing 20 mM Tris

buffer at a pH of 7.4, 4.2 mM MgCl_2 , and 1 mM dithiothreitol (DTT). Elution was performed with 50 mM Tris buffer at a pH of 9.6 containing 5 mM glutathione. The protein content in the fraction was determined by the pigment bonding method (Bradford method; BioRad Co.) and the main fraction was acquired.

Purity of the purified ribosomal protein L7/L12/GST fused protein that was obtained was confirmed by electrophoresis to be approximately 75 %, showing that a purity satisfactory for an immunogen could be guaranteed.

Example 3

Preparation of monoclonal antibody to ribosomal protein L7/L12 of *Haemophilus influenzae*

First, 100 μg fused protein antigen of ribosomal protein L7/L12/GST of *Haemophilus influenzae* were dissolved in 200 μl PBS and then 200 μl Freund's complete adjuvant were added and mixed and emulsification was performed. Two-hundred microliters were injected intraperitoneally to immunize mice.

Then the same emulsion antigen was intraperitoneally injected after 2 weeks, after 4 weeks, and after 6 weeks. Two-fold the concentration of antigen emulsion was injected intraperitoneally after 10 weeks and after 14 weeks. The spleen was

excised 3 days after the final immunization and submitted to cell fusion.

After thoroughly mixing 2×10^7 myeloma cells per 10^8 spleen cells from mice, which had been recovered
5 aseptically, in a glass tube, the mixture was centrifuged for 5 minutes at 1,500 rpm and the supernatant was discarded. The cells were thoroughly mixed.

The myeloma cells used for cell fusion were
10 obtained by cultivation of cell strain NS-1 with an RPMI 1640 culture medium containing 10 % FCS, cultivating this product beginning 2 weeks before cell fusion using an RPMI 1640 medium containing 0.13 mM azaguanine, 0.5 μ g/ml MC-210, and 10 % FCS for 1
15 weeks, and then further cultivating the cell strain for 1 week with an RPMI 1640 medium containing 10 % FCS.

Thirty milliliters of RPMI 1640 culture medium 50 ml that had been kept at 37 °C were added to the
20 mixed cell sample and centrifuged at 1,500 rpm. After removal of the supernatant, 1 ml 50 % polyethylene glycol that had been kept at 37 °C was added and stirred for 2 minute. 10 ml RPMI 1640 medium kept at 37 °C were added and the solution was
25 vigorously mixed for approximately 5 minutes as it was suctioned and evacuated from a sterile pipette.

After centrifugation for 5 minutes at 1,000 rpm

and removal of the supernatant, 30 ml HAT medium were added to bring the cell concentration to 5×10^6 cells/ml. This mixture was stirred till uniform and then poured, 0.1 ml at a time, into a 96-well culture plate and cultivated at 37 °C under 7 % CO₂. HAT medium was added, 0.1 ml at a time, on Day 1 and at Week 1 and Week 2.

Then the cells that had produced the desired antibody were screened by ELISA.

Solutions of GST fusion ribosomal protein L7/L12 and GST protein of *Haemophilus influenzae* dissolved in PBS containing 0.05 % sodium azide diluted to 10 µg/ml were poured, each 100 µl at a time, into separate 96-well plates and adsorbed overnight at 4 °C.

After removal of the supernatant, 200 µl of 1 % bovine serum albumin solution (in PBS) were added and reacted and blocked for 1 hour at room temperature. After removal of the supernatant, the product was washed with washing solution (0.02 % Tween 20, PBS). One-hundred microliters culture solution of fused cells were added to this and reacted for 2 hours at room temperature. The supernatant was removed and washed with washing solution. Next, 100 µl peroxidase-labeled goat anti-mouse IgG antibody solution at a concentration of 50 ng/ml were added and the solution was reacted for 1 hour at room

temperature. The supernatant was removed and the product was again washed with washing solution. Then TMB solution (KPL Co., ltd) was added, 100 μ l at a time, and the mixture was reacted for 20 minutes at room temperature. After coloration, 100 μ l 1 N sulfuric acid were added to stop the reaction and absorbance at 450 nm was determined.

As a result, positive wells that only reacted with GST fusion ribosomal protein L7/L12 and did not react with GST protein were detected, and it was concluded that antibody to ribosomal protein L7/L12 is present.

Therefore, the cells in the positive wells were recovered and cultivated with HAT medium in a 24-well plastic plate. The fused medium that had been cultivated was diluted with HT medium to a cell count of approximately 20 cells/ml and then mixed with 10^6 six-week-old mouse thyroid cells suspended in HT Cultivation medium in a 96-well culture plate. Culture was performed for 2 weeks at 37 °C under conditions of 7 % CO₂.

Antibody activity in the culture supernatant was similarly determined by the aforementioned ELISA method and the cells that showed positive reaction with ribosomal protein L7/L12 were recovered.

Furthermore, the same dilution test and cloning procedure was repeated to obtain a total of 5 clones

of hybridoma HIRB-1 ~ 5.

Example 4

Reaction of monoclonal antibody that reacts with
ribosomal protein L7/L12 of *Haemophilus influenzae*,
5 with *Neisseria gonorrhoeae* and other microorganisms

Monoclonal antibody was produced and recovered in accordance with standard methods using the positive hybridoma cells obtained as previously described.

Basically, 5×10^6 cells that had been
10 subcultured using RPMI 1640 culture medium
(containing 10 % FCS) were intraperitoneally injected
into Balb/C mice that had been intraperitoneally
injected with 0.5 ml Pristane 2 weeks earlier.
Ascites was recovered 3 weeks later and the
15 centrifugation supernatant was obtained.

The solution containing antibody that was
obtained was adsorbed in a Protein A column (5 ml
bed, Pharmacia) and washed with 3-bed volume of PBS.
Then eluted with citrate buffer at pH 3, the antibody
20 fraction was recovered and the monoclonal antibody
produced by each hybridoma was obtained.

The monoclonal antibody derived from these 5
strains of hybridoma was used in ELISA.

The sandwich assay method was used to assess the
25 monoclonal antibody. The monoclonal antibody that
was prepared was used as an enzyme-labeled antibody
by being chemically bound to peroxidase.

That is, enzyme labeling was performed in accordance with the method in "Analytical Biochemistry" 132 (1983), 68-73 with the reagent S-acetylthioacetic acid N-hydroxysuccinimide for
5 binding using horseradish peroxidase (Sigma Grade VI). By means of the ELISA reaction, a solution of commercial anti-*Haemophilus influenzae* polyclonal antibody which dissolved in PBS containing 0.05 % sodium azide, (Biodesign, rabbit) was diluted to a
10 concentration of 10 µg/ml and poured, 100 µl at a time, into a separate 96-well plate and adsorbed overnight at 4 °C.

After removal of the supernatant, 200 µl 1 % FCS solution (in PBS) were added and reacted and blocked
15 for 1 hour at room temperature. The supernatant was removed and the product was washed with washing solution (0.02 % Tween 20, PBS). One-hundred microliters of antigen solution, which had been obtained by adding Triton X-100 to culture solutions
20 of each species of microorganism to a concentration of 0.3 % and then extracting the solution for 5 minutes at room temperature, were added to this and reacted for 2 hours at room temperature. The supernatant was removed and the product was further
25 washed with washing solution. Then 100 µl peroxidase-labeled anti-ribosomal protein L7/L12 antibody solution at 5 µg/ml were added and reacted

for 1 hour at room temperature. The supernatant was removed and the product was washed with washing solution. TMB (KPL Co.) solution was added, 100 μ l at a time, and reacted for 20 minutes at room

5 temperature. After coloration, 100 μ l 1 N sulfuric acid were added to stop the reaction. Absorbance at 450 nm was determined.

As a result, as shown in Table 1 it is clear that when monoclonal antibody derived from hybridoma HIRB-
10 2 was used as the enzyme-labeled antibody, all strains of *Haemophilus influenzae* tested were detected at a sensitivity of 10^6 bacteria/ml, while reactivity of other bacteria belonging to the genus *Neisseria* and other microorganisms could not be
15 detected, even at high concentrations of 10^8 bacteria/ml and therefore, antibody with specific reactivity to *Haemophilus influenzae* can be obtained by using monoclonal antibody to ribosomal protein L7/L12.

20

25

Table 1

Results of Detection (10 ⁶ cells/ml)		
<i>Haemophilus influenzae</i>	ATCC9327	+
<i>Haemophilus influenzae</i>	ATCC9334	+
<i>Haemophilus influenzae</i>	ATCC9007	+
<i>Haemophilus influenzae</i>	ATCC9332	+
<i>Haemophilus influenzae</i>	ATCC8142	+
<i>Haemophilus influenzae</i>	ATCC9833	+
Results of Detection (10 ⁸ cells/ml)		
<i>Neisseria meningitides</i>	ATCC13090	—
<i>Neisseria lactamica</i>	ATCC30011	—
<i>Neisseria mucosa</i>	ATCC35611	—
<i>Neisseria sicca</i>	ATCC9913	—
<i>Branhamella catarrharis</i>	ATCC25240	—
<i>Neisseria gonorrhoeae</i>	ATCC9793	—
<i>Escherichia coli</i>	ATCC25922	—
<i>Klebsiella pneumoniae</i>	ATCC13883	—

5 (+: Positive; -: Negative)

Example 5

10 Cloning of ribosomal protein L7/L12 genes from
Streptococcus pneumponiae, mass expression in
Escherichia coli and purification of the same protein
and preparation of monoclonal antibody to the same
protein

After inoculating an appropriate amount of
Streptococcus pneumoniae strain IID555 (obtained from

Tokyo University School of Medicine Laboratories) in a blood agar culture medium, the strain was cultivated for 48 hours in an incubator at 37 °C. The colonies that grew were suspended in a TE buffer to a
5 final concentration of approximately 5×10^9 CFU/ml. Approximately 1.5 ml of this suspension was transferred to a microcentrifuge tube and centrifuged for 2 minutes at 10,000 rpm. The supernatant was discarded. The sediment was resuspended in 567 µl TE
10 buffer. Then 30 µl 10 % SDS and 3 µl 20 mg/ml Proteinase K solution were added and thoroughly mixed. The suspension was incubated for another hour at 37 °C. Next, after adding 80 µl 10 % cetyl trimethyl ammonium bromide/0.7 M NaCl solution and
15 thoroughly mixing the product, it was incubated for 10 minutes at 65 °C. Next, 700 µl chloroform-isoamyl alcohol solution at a volume ratio of 24:1 was added and stirred well. The solution was centrifuged for 5 minutes (while being kept at 4 °C) at 12,000 rpm
20 using a microcentrifugation device and the aqueous fraction was transferred to a new microtube. Isopropanol was added to the fraction at 0.6-times its volume and the tube was vigorously shaken to form sediment of the DNA. The white DNA sediment was
25 scooped with a glass rod and transferred to a different microcentrifuge tube containing 1 ml 70 % ethanol (cooled to -20 °C).

Next, the product was centrifuged for 5 minutes at 10,000 rpm and the supernatant was gently removed. Then another 1 ml 70 % ethanol was added and the product was centrifuged for 5 more minutes. Once the supernatant had been removed, the sediment was dissolved in 100 μ l TE buffer to obtain the DNA solution. The concentration of the genomic DNA solution was determined quantitatively in accordance with E5, Spectrophotometric determination of the amount of DNA or RNA, "Molecular cloning: A laboratory manual," 1989, Eds. Sambrook, J., Fritsch, E.F., and Maniatis, T., Cold Spring Harbor Laboratory Press.

1105
13
PCR was performed using 10 ng of this genomic DNA. Taq polymerase (Takara Co., ltd., code R001A) was employed for PCR. Then 5 μ l of buffer attached to enzyme, 4 μ l dNTP mixture attached to enzyme, and 200 pmol each of synthetic oligonucleotide C shown in Sequence No. 13 of the Sequence Table and synthetic oligonucleotide D shown in Sequence No. 14 of the Sequence table were added to the enzyme to bring the final volume to 50 μ l.

This mixture was cycled 5 times with a TaKaRa PCR Thermal Cycler 480 for 1 minute at 95 $^{\circ}$ C, 2 minutes at 50 $^{\circ}$ C, and 3 minutes at 72 $^{\circ}$ C and was then cycled 25 times for 1 minute at 95 $^{\circ}$ C, 2 minutes at 60 $^{\circ}$ C, and 3 minutes at 72 $^{\circ}$ C. Electrophoresis was performed

in 1.5 % agarose gel using some of this PCR product.
This product was then stained with ethidium bromide
(Nihon Gene Co., ltd.) and observed under ultraviolet
rays to confirm amplification of approximately 400 bp
5 cDNA. After digestion treatment with restriction
endonucleases BamHI and XhoI, electrophoresis was
performed in 1.5 % agarose gel and staining with
ethidium bromide was carried out. An approximately
370 bp band was cut out from the gel. This band was
10 purified with Suprec01 (Takara Co., Ltd.) and then
inserted into pGEX-6P-1 (Pharmacia), which is a
commercial vector.

This same vector can function as an expression
vector for the desired molecule, which can express
15 fused protein with GST protein, by insertion of the
desired gene fragment into the appropriate
restriction endonuclease site. Actually, vector
pGEX-6P-1 and the previous DNA were mixed together at
a molar ratio of 1:5 and DNA was inserted into the
20 vector with T4 DNA ligase (Invitrogen Co.). Vector
pGEX-4T-1 into which DNA had been inserted was
genetically introduced to *Escherichia coli* One-Shot
Competent Cells (Invitrogen Co., Ltd.) and then
inoculated in a plate of L-Broth (Takara Co., Ltd.)
25 semi-solid culture plate containing 50 µg/ml
ampicillin (Sigma). The plate was then set aside at
37°C for 12 hours and the colonies that grew were

selected at random and inoculated into 2 ml L-Broth liquid culture medium containing the same concentration of ampicillin. Shake cultivation was performed at 37°C for 8 hours and the bacteria was recovered and the plasmid was separated using Wizard Miniprep (Promega Co.,) in accordance with the attached literature. The plasmid was cleaved with restriction endonuclease BamHI/XhoI. Insertion of said PCR product was confirmed by cutting out approximately 370 bp DNA. The base sequence of the DNA that had been inserted was determined using said clone.

Determination of the base sequence of the inserted DNA fragment was performed using the Fluorescence Sequencer of Applied Biosystems. The sequence sample was prepared using PRISM, Ready Reaction Dye Terminator Cycle Sequencing Kit (Applied Biosystems). First, 9.5 µl reaction stock solution, 4.0 µl T7 promoter primer at 0.8 pmol/µl (Gibco BRL) and 6.5 µl of template DNA for sequencing at 0.16 µg/µl were added to a microtube with a capacity of 0.5 ml, mixed and superposed with 100 µl mineral oil. PCR amplification was performed for 25 cycles, where one cycle consisted of 30 seconds at 96 °C, 15 seconds at 55 °C, and 4 minutes at 60 °C. The product was then kept at 4 °C for 5 minutes. After the reaction was completed, 80 µl sterilized pure water

was added and stirred. The product was centrifuged and the aqueous layer was extracted 3 times with phenol-chloroform. Ten microliters 3 M sodium acetate (pH 5.2) and 300 μ l ethanol were added to 100 μ l aqueous layer and stirred. The product was then centrifuged for 15 minutes at room temperature and 14,000 rpm and the sediment was recovered. Once the sediment was washed with 75 % ethanol, it was dried under a vacuum for 2 minutes to obtain the sequencing sample. The sequencing sample was dissolved in formamide containing 4 μ l 10 mM EDTA and denatured for 2 minutes at 90 °C. This was then cooled in ice and submitted to sequencing.

One of the 7 clones obtained had homology of the sequence with the probe used for PCR. In addition, DNA sequences extremely similar to the gene sequence of ribosomal protein L7/L12 gene of the other microorganisms, for example, *Neisseria gonorrhoeae*, were discovered. The entire base sequence and the corresponding amino acid sequence of the structural gene moiety are as shown in Sequence ID No. 19 and No. 20 of the Sequence Table. This gene fragment clearly codes for *Streptococcus pneumoniae* ribosomal protein L7/L12.

50 ml *Escherichia coli* into which expression vector had been inserted was cultivated overnight in a two-fold concentration YT medium at 37 °C. Then,

450 ml of the two-fold concentration YT medium was heated at 37 °C for 1 hour. 50 ml of the *Escherichia coli* solution that had been cultivated overnight was introduced to 450 ml of the aforementioned medium.

- 5 After cultivation for one hour at 37 °C, 100 µl 500 mM IPTG was introduced and cultivated for 4 hours at 25 °C. The product was then recovered and introduced to centrifugation tube at each 250 ml and centrifuged for 20 minutes at 5000 rpm. The supernatant was
10 discarded and dissolved in 25 ml each 50 mM Tris-HCl at a pH of 7.4 and Lysis buffer containing 25 % sucrose.

- Furthermore, 1.25 ml 10 % NP-40 and 125 µl 1 M MgCl₂ were added and transferred to a plastic tube.
15 Sonication was performed 1 minute x 5 times while ice cold. The product was centrifuged for 15 minutes at 12,000 rpm and the supernatant was recovered.

- Next, the aforementioned supernatant was adsorbed on a glutathione sepharose column (manufactured by
20 Pharmacia) conditioned with PBS. Then, the column was washed with PBS three times the bed volume. Elution was performed with 50 mM Tris-HCl at a pH of 8.0 containing 10 mM glutathione. The protein content in the fraction was determined by the pigment
25 bonding method (Bradford method; BioRad Co.) and the main fraction was acquired. The main fraction was dialyzed three times against 3 L PBS.

1 ml of a cleavage buffer containing 500 mM Tris-HCl (pH 7.0), 1.5 M NaCl, 10 mM EDTA, and 10 mM DTT was added to 10 ml of 1 mg/ml solution of the resulting GST fusion ribosomal protein L7/L12. 100 μ l of 2 u/ μ l PreScission Protease (manufactured by Pharmacia company) was further added and reacted at 4 °C to separate the GST moiety from ribosomal protein L7/L12.

The reaction solution was passed through a glutathione sepharose column which had been conditioned with PBS. The solution coming out from the column was recovered. One-bed volume of PBS was passed through and also recovered. Purity of the purified ribosomal protein L7/L12 that was obtained was confirmed by electrophoresis to be approximately 90 %, showing that a purity satisfactory for an immunogen could be guaranteed.

First, 100 μ g protein antigen of ribosomal protein L7/L12 of *Streptococcus pneumoniae* was dissolved in 200 μ l PBS and then 200 μ l Freund's complete adjuvant was added and mixed and emulsification was performed. 200 μ l was intraperitoneally injected to immunize mice. Then, the same emulsion antigen was intraperitoneally injected after 2 weeks, after 4 weeks, and after 6 weeks. A two-fold concentration antigen emulsion was further injected intraperitoneally after 10 weeks and

after 14 weeks. The spleen was excised 3 days after the final immunization and submitted to cell fusion.

After thoroughly mixing 2×10^7 myeloma cells per 10^8 spleen cells from mice, which had been recovered
5 aseptically, in a glass tube, the mixture was centrifuged for 5 minutes at 1500 rpm and the supernatant was discarded. The cells were thoroughly mixed.

The myeloma cells used for cell fusion were
10 obtained by cultivation of cell strain NS-1 with an RPMI 1640 culture medium containing 10 % FCS, cultivating this product beginning 2 weeks before cell fusion using an RPMI 1640 medium containing 0.13 mM azaguanine, 0.5 μ g/ml MC-210, and 10 % FCS for 1
15 weeks, and then further cultivating the cell strain for 1 week with an RPMI 1640 medium containing 10 % FCS. 30 ml of RPMI 1640 culture medium 50 ml that had been kept at 37 °C was added to the mixed cell sample and centrifuged at 1,500 rpm. After removal of the
20 supernatant, 1 ml 50 % polyethylene glycol that had been kept at 37 °C was added and stirred for 2 minute. 10 ml RPMI 1640 medium kept at 37 °C was added and the solution was vigorously mixed for approximately 5 minutes as it was suctioned and
25 evacuated using a sterile pipette.

After centrifugation for 5 minutes at 1,000 rpm and removal of the supernatant, 30 ml HAT medium were

added to bring the cell concentration to 5×10^6 cells/ml. This mixture was stirred till uniform and then poured, 0,1ml at a time, into a 96-well culture plate and cultivated at 37 °C under 7 % CO₂. HAT
5 medium was added, 0.1 ml at a time, on Day 1 and at Week 1 and Week 2.

Then, the cells that had produced the desired antibody were screened by ELISA. Solutions of ribosomal protein L7/L12 of *Streptococcus pneumoniae*
10 dissolved in PBS containing 0.05 % sodium azide diluted to 10 µg/ml were poured, 100 µl at a time, into separate 96-well plates and adsorbed overnight at 4 °C. After removal of the supernatant, 200 µl 1 % bovine serum albumin solution (in PBS) were added and
15 reacted and blocked for 1 hour at room temperature. The supernatant was removed and the product was washed with a washing solution (0.02 % Tween 20, PBS). 100 µl of a culture solution of fusion cells was added and reacted for two hours at room
20 temperature. The supernatant was removed and the product was further washed with a washing solution. Then, 100 µl of a peroxidase-labeled goat anti-mouse antibody solution at 50 ng/ml was added and reacted for one hour at room temperature. The supernatant
25 was removed and the product was washed with a washing solution. TMB (KPL) solution was added, 100 µl at a time, and reacted for 20 minutes at room temperature.

After coloration, 100 μ l 1 N sulfuric acid were added to stop the reaction. Absorbance at 450 nm was determined.

As a result, positive wells that reacted with
5 ribosomal protein L7/L12 were detected, confirming presence of the antibody to ribosomal protein L7/L12.

Therefore, the cells in the positive wells were recovered and cultivated with HAT medium in a 24-well plastic plate. The fused medium that had been
10 cultivated was diluted with an HT medium to a cell count of approximately 20 cells/ml and then mixed with 10^6 six-week-old mouse thyroid cells suspended in the HT medium in a 96-well culture plate. The cells were cultivated for 2 weeks at 37 °C under the
15 conditions of 7 % CO₂. The antibody activity in the culture supernatant was determined by the aforementioned ELISA method and the cells that showed a positive reaction with ribosomal protein L7/L12 were recovered.

20 Furthermore, the same dilution and cloning procedure was repeated to obtain a total of 4 clones of hybridoma AMSP-1 to 4.

Example 6

Reaction of monoclonal antibody that reacts with
25 ribosomal protein L7/L12 of *Streptococcus pneumoniae*,
with *Streptococcus pneumoniae* and other
microorganisms

A monoclonal antibody was produced and recovered in accordance with standard methods using the positive hybridoma cells obtained as previously described.

5 Specifically, cells subcultured in RPMI 1640 medium (containing 10 % FCS) was diluted with a serum-free medium to about 2×10^5 cells/ml, 3.3×10^5 cells/ml, and 5×10^5 cells/ml in 25 cm² culture flasks, and the total volume was made 5 ml. After
10 cells were grown for 3 to 5 days in 7 % CO₂ at 37 °C, a flask which contains the least number of original cells was selected among flasks in which cells were grown. The same procedure was repeated until the cells diluted to 2×10^5 cells/ml grow to 2×10^6
15 cells/ml in 3 to 4 days, thereby adapting the cells with the serum-free medium. Next, cloning was performed in a 96-well plate for bacteria cultivation to select cells exhibiting fastest growth and a highest antibody titer. The selected cells were
20 grown in a 24-well plate and diluted with a serum-free medium in a 25 cm² culture flask to a concentration of about 2×10^5 cells/ml and the total volume was made 10 ml. After incubation for 3 to 4 days in 7 % CO₂ at 37 °C to a concentration of 1×10^6
25 cells/ml, the culture broth 100 ml, 1×10^6 cells/ml was transferred to a bottle for mass cultivation which were grown in the same manner in a 75 cm²

flask. 100 ml of a serum-free medium was added to the mixture, which was incubated at 37 °C for two days while stirring. 200 ml of the serum-free medium was added again and the mixture was incubated for a further two days. The culture broth was divided into four aliquot, the serum-free medium was added to each portion, followed by incubation for two days. After further addition of 400 ml of the serum-free medium, the culture broth was incubated for 6 days. The culture broth was collected and centrifuged at 10,000 rpm for 15 minutes to obtain a culture supernatant that contain the target antibody. After the addition of sodium azide to final concentration 0.1 %, the culture supernatant was stored at 4 °C. 100 ml of the solution containing the antibody that was obtained was 5-fold diluted with PBS and adsorbed in a Protein G column (5 ml bed, Pharmacia) and washed with 3-bed volume of PBS. Then eluted with citrate buffer at a pH 3, the antibody fraction was recovered, and monoclonal antibody produced by each hybridoma was obtained. The monoclonal antibodies originating from the four hybridoma clones were evaluated according to the OIA method described in Publication of International Patent Application Japanese Laid-open (Toku-Hyou) No. 509565/1995.

Specifically, the OIA method comprises preparing a reactive substrate by reacting an antibody for

capture on a silicon wafer having a thin film layer of silicon nitride, causing this substrate to react with an antigen which is an extract of microorganisms for a prescribed period of time, causing the captured antigen to react with an antibody (an amplification reagent) which is an enzyme-labeled antibody, and finally adding a substrate solution to produce a thin-film precipitate. The antigen-antibody reaction can be judged visually by a degree of light interference color produced in the precipitate.

The monoclonal antibody preparation was used and evaluated as a capture antibody to be immobilized on a silicon wafer having a silicon nitride thin film layer in the OIA method. Moreover, peroxidase-labeled AMGC-1 monoclonal antibody which can non-specifically react with ribosomal proteins L7/L12 protein of a variety of microorganisms described in Reference Example was used as the detect antibody. That is, enzyme labeling was performed in accordance with the method in "Analytical Biochemistry" 132 (1983), 68-73 with the reagent S-acetylthioacetic acid N-hydroxysuccinimide for binding using horseradish peroxidase (Sigma Grade VI).

In the OIA reaction, monoclonal antibody in a PBS containing 0.05 % sodium azide was diluted with 0.1 M HEPES (pH 8.0) to a concentration of 10 µg/ml and added onto a silicone wafer which has a thin film

layer of silicon nitride, 50 μ l at a time, to react for 30 minutes at room temperature, followed by washing with distilled water and use.

15 μ l of antigen solution, which had been
5 obtained by adding 0.5% Triton X-100 to culture solutions of various species of microorganisms and then extracting the solution for 5 minutes at room temperature, was added to the specimen obtained in the above-described procedure and reacted for 10
10 minutes at room temperature. Then, 15 μ l of 20 μ g/ml peroxidase-labeled AMGC1 anti-body was added and reacted for 10 minutes. After washing with distilled water, a substrate solution (KPL) was added, 15 μ l at a time, and reacted for 5 minutes at room
15 temperature. The product was washed with distilled water to judge the concentration of detection signals as an intensity of light interference by naked eyes.

As a result, as shown in Table 2 it is clear that when monoclonal antibody derived from hybridoma AMSP-
20 2 was used as the capture antibody, all strains of *Streptococcus pneumoniae* tested were detected at a sensitivity of 10^6 bacteria/ml, while reactivity of other bacteria could not be detected at a higher concentration of 10^8 bacteria/ml. Thus, the antibody
25 with specific reactivity to *Streptococcus pneumoniae* was confirmed to have been obtained by using the monoclonal antibody to ribosomal protein L7/L12.

The hybridoma AMSP-2 which produces the monoclonal antibody specific to *Streptococcus pneumoniae* has been deposited with National Institute of Bioscience and Human-Technology, the Agency of Industrial Science and Technology, the Ministry of International Trade and Industry, Japan, on July 28, 1999, with the disposition number FERM BP-6807.

Table 2

Results of Detection (10^6 cells/ml)		
<i>Streptococcus pneumoniae</i>	ATCC27336	+
<i>Streptococcus pneumoniae</i>	IID554	+
<i>Streptococcus pneumoniae</i>	IID555	+
<i>Streptococcus pneumoniae</i>	IID556	+
<i>Streptococcus pneumoniae</i>	IID557	+
<i>Streptococcus pneumoniae</i>	IID558	+
<i>Streptococcus pneumoniae</i>	IID559	+
<i>Streptococcus pneumoniae</i>	IID1603	+
Results of Detection (10^8 cells/ml)		
<i>Escherichia coli</i>	ATCC25922	-
<i>Enterococcus faecalis</i>	ATCC19433	-
<i>Haemophilus influenzae</i>	ATCC10211	-
<i>Klebsiella pneumoniae</i>	ATCC13883	-
<i>Neisseria gonorrhoeae</i>	IID821	-
<i>Neisseria lactamica</i>	ATCC23970	-
<i>Neisseria meningitides</i>	ATCC13090	-
<i>Pseudomonas aeruginosa</i>	ATCC27853	-
Group B streptococcus	ATCC12386	-
<i>Staphylococcus aureus</i>	ATCC25923	-
<i>Streptococcus pyogenes</i>	ATCC19615	-

(+: Positive; -: Negative)

Example 7

Cloning of ribosomal protein L7/L12 genes from
Neisseria gonorrhoeae, mass expression in *Escherichia*
coli and purification of the same protein and
5 preparation of monoclonal antibody to the same
protein

1WS
B14
10 After inoculating an appropriate amount of
Neisseria gonorrhoeae strain IID821 (obtained from
Tokyo University School of Medicine Laboratories) in
a chocolate agar culture medium, the strain was
cultivated for 24 hours in a CO₂ incubator under
conditions of 37 °C and 0.5 <5.0> % CO₂. The colonies
that grew were suspended in a TE buffer to a final
concentration of approximately 5 x 10⁹ CFU/ml.
15 Approximately 1.5 ml of this suspension was
transferred to a microcentrifugation tube and
centrifuged for 2 minutes at 10,000 rpm. The
supernatant was discarded. The sediment was
resuspended in 567 µl TE buffer. Then 30 µl 10 % SDS
20 and 3 µl 20 mg/ml Proteinase K solution were added
and thoroughly mixed. The suspension was incubated
for another hour at 37 °C.

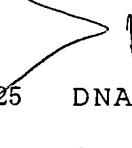
Next, after adding 80 µl 10 % cetyl trimethyl
ammonium bromide/0.7 M NaCl solution and thoroughly
25 mixing the product, it was incubated for 10 minutes
at 65 °C. Next, 700 µl chloroform-isoamyl alcohol
solution at a volume ratio of 24:1 was added and

stirred well. The solution was centrifuged for 5 minutes (while being kept at 4 °C) at 12,000 rpm using a microcentrifugation device and the aqueous fraction was transferred to a new microtube.

- 5 Isopropanol was added to the fraction at 0.6-times its volume and the tube was vigorously shaken to form sediment of the DNA. The white DNA sediment was scooped with a glass rod and transferred to a different microcentrifugation tube containing 1 ml 70 % ethanol (cooled to -20 °C).

Next, the product was centrifuged for 5 minutes at 10,000 rpm and the supernatant was gently removed. Then another 1 ml 70 % ethanol was added and the product was centrifuged for 5 more minutes.

- 15 Once the supernatant had been removed, the sediment was dissolved in 100 µl TE buffer to obtain the DNA solution. The concentration of the genomic DNA solution was determined quantitatively in accordance with E5, Spectrophotometric determination of the amount of DNA or RNA, "Molecular cloning: A laboratory manual," 1989, Eds. Sambrook, J., Fritsch, E.F., and Maniatis, T., Cold Spring Harbor Laboratory Press.

- 25 *INS*
BIS  PCR was performed using 10 ng of this genomic DNA. PCR was performed using Taq polymerase (Takara Co., Ltd., code R001A). Then, 5 µl of a buffer attached to enzyme, 4 µl of a dNTP mixture attached

to enzyme, and 200 pmol each of synthetic oligonucleotide E shown in Sequence No. 15 of the Sequence Table and synthetic oligonucleotide F shown in Sequence No. 16 of the Sequence Table, which were
5 designed based on the ribosomal protein L7/L12 DNA sequence of *Neisseria gonorrhoeae* acquired from Internet Information (Oklahoma University, N. *Gonorrhoeae* Genome Project, disclosed genomic DNA data) because of the similarity with ribosomal
10 protein L7/L12 DNA sequence of other bacteria, were added to the enzyme to bring the final volume to 50 μ l.

This mixture was cycled 5 times with a TaKaRa PCR Thermal Cycler 480 for 1 minute at 95 °C, 2 minutes
15 at 50 °C, and 3 minutes at 72 °C and was then cycled 25 times for 1 minute at 95 °C, 2 minutes at 60 °C, and 3 minutes at 72 °C. Electrophoresis was performed in 1.5 % agarose gel using some of this PCR product. This product was then stained with ethidium bromide
20 (Nihon Gene Co., ltd.) and observed under ultraviolet rays to confirm amplification of approximately 400 bp cDNA. After digestion treatment with restriction endonucleases BamHI and XhoI, electrophoresis was performed in 1.5 % agarose gel and staining with
25 ethidium bromide was carried out. An approximately 370 bp band was cut out from the gel. This band was purified with Suprec01 (Takara Co., Ltd.) and then

inserted into pGEX-4T-1 (Pharmacia), which is a commercial vector. Actually, vector pGEX-4T-1 and the previous DNA were mixed together at a molar ratio of 1:3 and DNA was inserted into the vector with T4 DNA ligase (Invitrogen Co.). Vector pGEX-4T-1 into which DNA had been inserted was genetically introduced to *Escherichia coli* One-Shot Competent Cells (Invitrogen Co., Ltd.) and then inoculated in a plate of L-Broth (Takara Co., Ltd.) semi-solid culture plate containing 50 µg/ml ampicillin (Sigma). The plate was then set aside at 37 °C for 12 hours and the colonies that grew were selected at random and inoculated into 2ml L-Broth liquid culture medium containing the same concentration of ampicillin. Shake cultivation was performed at 37°C for 8 hours and the bacteria was recovered and the plasmid was separated using Wizard Miniprep in accordance with the attached literature. The plasmid was cleaved with restriction endonuclease BamHI/XhoI. Insertion of said PCR product was confirmed by cutting out approximately 370 bp DNA. The base sequence of the DNA that had been inserted was determined using said clone.

INS
B16
Determination of the base sequence of the inserted DNA fragment was performed using the Fluorescence Sequencer of Applied Biosystems. The sequence sample was prepared using PRISM, Ready Reaction Dye Terminator Cycle Sequencing Kit (Applied

Biosystems). First, 9.5 μ l reaction stock solution, 4.0 μ l T7 promoter primer at 0.8 pmol/ μ l (Gibco BRL) and 6.5 μ l template DNA for sequencing at 0.16 μ g/ μ l were added to a microtube with a capacity of 0.5 ml and mixed. After layering with 100 μ l mineral oil, PCR amplification was performed for 25 cycles, where one cycle consisted of 30 seconds at 96 °C, 15 seconds at 55 °C, and 4 minutes at 60 °C. The product was then kept at 4 °C for 5 minutes. After the reaction was completed, 80 μ l sterilized pure water was added and stirred. The product was centrifuged and the aqueous layer was extracted 3 times with phenol-chloroform. Ten microliters 3 M sodium acetate (pH 5.2) and 300 μ l ethanol were added to 100 μ l aqueous layer and stirred. The product was then centrifuged for 15 minutes at room temperature and 14000 rpm and the sediment was recovered. Once the sediment was washed with 75 % ethanol, it was dried under a vacuum for 2 minutes to obtain the sequencing sample. The sequencing sample was dissolved in formamide containing 4 μ l 10 mM EDTA and denatured for 2 minutes at 90 °C. This was then cooled in ice and submitted to sequencing. One of the 5 clones obtained had homology of the sequence with the probe used for PCR. In addition, DNA sequences extremely similar to the gene sequence of ribosomal protein L7/L12 gene of the other microorganisms, for example,

Haemophilus influenzae, were discovered. The entire base sequence and the corresponding amino acid sequence of the structural gene moiety are as shown in Sequence No. 21 and No. 22 of the Sequence Table.

5 This gene fragment clearly codes for ribosomal protein L7/L12 gene of *Neisseria gonorrhoeae*.

Neisseria gonorrhoeae fusion GST ribosome protein L7/L12 prepared by the same method as in Example 2 was obtained using the *Neisseria gonorrhoeae* fusion
10 GST ribosomal protein L7/L12 expression vector constructed in this way.

Furthermore, hybridoma strain GCRB-3, which produces monoclonal antibody to ribosomal protein L7/L12 of *Neisseria gonorrhoeae*, was obtained in
15 accordance with the method in Example 3.

Example 8

Reaction of monoclonal antibody that reacts with ribosomal protein L7/L12 of *Neisseria gonorrhoeae*, with *Neisseria gonorrhoeae* and other microorganisms

20 Monoclonal antibody was produced and recovered in accordance with standard methods using the positive hybridoma cells GCRB-3 obtained as previously described.

Basically, 5×10^6 cells (in PBS) that had been
25 subcultured using RPMI 1640 culture medium (containing 10 % FCS) were intraperitoneally injected into Balb/C mice that had been intraperitoneally

injected with 0.5 ml Pristane 2 weeks earlier.
Ascites was recovered 3 weeks later and the
centrifugation supernatant was obtained.

The solution containing antibody that was
5 obtained was adsorbed in a Protein A column (5 ml
bed, Pharmacia) and washed with 3-bed volume of PBS.
Then eluted with citrate buffer at pH 3, the antibody
fraction was recovered and the monoclonal antibody
that was produced by each hybridoma was obtained.
10 The monoclonal antibody derived from the GCRB-3
hybridoma was used in ELISA.

The sandwich assay method was used to assess the
monoclonal antibody. The monoclonal antibody that
was prepared was used as an enzyme-labeled antibody
15 by being chemically bound to peroxidase. That is,
enzyme labeling was performed in accordance with the
method in "Analytical Biochemistry" 132 (1983), 68-73
with the reagent S-acetylthioacetic acid N-
hydroxysuccinimide for binding using horseradish
20 peroxidase (Sigma Grade VI). By means of the ELISA
reaction, a solution of commercial anti-*Neisseria*
gonorrhoeae polyclonal antibody in PBS containing
0.05% sodium azide (Virostat, rabbit) was diluted to a
concentration of 10 µg/ml and poured, 100 µl at a
25 time, into a separate 96-well plate and adsorbed
overnight at 4 °C.

After removal of the supernatant, 200 µl 1 %

bovine serum albumin solution (in PBS) were added and reacted for 1 hour and blocked at room temperature. The supernatant was removed and the product was washed with washing solution (0.02 % Tween 20, PBS).

- 5 One-hundred microliters of antigen solution, which had been obtained by adding Triton X-100 to culture solutions of each species of microorganisms to a concentration of 0.3 % and then extracting the solution for 5 minutes at room temperature, were
- 10 added to this and reacted for 2 hours at room temperature. The supernatant was removed and the product was further washed with washing solution. Then 100 μ l peroxidase-labeled anti-ribosomal protein L7/L12 antibody solution at 5 μ g/ml were added and
- 15 reacted for 1 hour at room temperature. The supernatant was removed and the product was washed with washing solution. TMB (KPL) solution was added, 100 μ l at a time, and reacted for 20 minutes at room temperature. After coloration, 100 μ l 1 N sulfuric
- 20 acid were added to stop the reaction. Absorbance at 450 nm was determined.

As a result, as shown in Table 3 it is clear that when monoclonal antibody derived from hybridoma GCRB-3 was used as the enzyme-labeled antibody, all

25 strains of *Neisseria gonorrhoeae* tested were detected at a sensitivity of 10^6 cells/ml, while reactivity of other species belonging to the genus *Neisseria* and

other microorganisms could not be detected, even at high concentrations of 10^8 cells/ml and therefore, antibody with specific reactivity to *Neisseria gonorrhoeae* can be obtained by using monoclonal antibody to ribosomal protein L7/L12.

Table 3

Results of Detection (10^6 cells/ml)		
<i>Neisseria gonorrhoeae</i>	ATCC9793	+
<i>Neisseria gonorrhoeae</i>	ATCC19424	+
<i>Neisseria gonorrhoeae</i>	ATCC27628	+
<i>Neisseria gonorrhoeae</i>	ATCC27629	+
<i>Neisseria gonorrhoeae</i>	ATCC27630	+
<i>Neisseria gonorrhoeae</i>	ATCC27631	+
<i>Neisseria gonorrhoeae</i>	ATCC27632	+
<i>Neisseria gonorrhoeae</i>	ATCC27633	+
<i>Neisseria gonorrhoeae</i>	ATCC35541	+
<i>Neisseria gonorrhoeae</i>	ATCC35542	+
<i>Neisseria gonorrhoeae</i>	ATCC43069	+
<i>Neisseria gonorrhoeae</i>	ATCC43070	+
<i>Neisseria gonorrhoeae</i>	ATCC49226	+
Results of Detection (10^8 cells/ml)		
<i>Neisseria meningitidis</i>	ATCC13090	—
<i>Neisseria lactamica</i>	ATCC30011	—
<i>Neisseria mucosa</i>	ATCC35611	—
<i>Neisseria sicca</i>	ATCC9913	—
<i>Branhamella catarrharis</i>	ATCC25240	—
<i>Haemophilus influenzae</i>	ATCC10211	—
<i>Escherichia coli</i>	ATCC25922	—
<i>Klebsiella pneumoniae</i>	ATCC13883	—

10 (+: Positive; -: Negative).

Example 9

Acquisition of ant-ribosomal protein L7/L12
monoclonal antibody specific to genus *Neisseria*

5 After inoculating an appropriate amount of
Neisseria gonorrhoeae strain IID821 (obtained from
Tokyo University School of Medicine Laboratories) in
a chocolate agar culture medium, the strain was
cultivated for 24 hours in a CO₂ incubator under
10 conditions of 37 °C and 0.5 <5.0> % CO₂. The colonies
that grew were suspended in a TE buffer to a final
concentration of approximately 5 x 10⁹ CFU/ml.
Approximately 1.5 ml of this suspension was
transferred to a microcentrifugation tube and
15 centrifuged for 2 minutes at 10,000 rpm. The
supernatant was discarded. The sediment was
resuspended in 567 µl TE buffer. Then 30 µl 10 % SDS
and 3 µl 20 mg/ml Proteinase K solution were added
and thoroughly mixed. The suspension was incubated
20 for another hour at 37 °C. Next, after adding 80 µl
10 % cetyl trimethyl ammonium bromide/0.7 M NaCl
solution and thoroughly mixing the product, it was
incubated for 10 minutes at 65 °C. Next, 700 µl
chloroform-isoamyl alcohol solution at a volume ratio
25 of 24:1 was added and stirred well.

The solution was centrifuged for 5 minutes (while
being kept at 4 °C) at 12,000 rpm using a

microcentrifugation device and the aqueous fraction was transferred to a new microtube. Isopropanol was added to the fraction at 0.6-times its volume and the tube was vigorously shaken to form sediment of the DNA. The white DNA sediment was scooped with a glass rod and transferred to a different microcentrifugation tube containing 1 ml 70 % ethanol (cooled to -20 °C).

Next, the product was centrifuged for 5 minutes at 10,000 rpm and the supernatant was gently removed. Then another 1 ml 70 % ethanol was added and the product was centrifuged for 5 more minutes. Once the supernatant had been removed, the sediment was dissolved in 100 µl TE buffer to obtain the DNA solution. The concentration of the genomic DNA solution was determined quantitatively in accordance with E5, Spectrophotometric determination of the amount of DNA or RNA, "Molecular cloning: A laboratory manual," 1989, Eds. Sambrook, J., Fritsch, E.F., and Maniatis, T., Cold Spring Harbor Laboratory Press.

PCR was performed using 10 ng of this genomic DNA. Taq polymerase (Takara Co., Ltd., code R001A) was employed for PCR. Then, 5 µl buffer attached to enzyme, 4 µl dNTP mixture attached to enzyme, and 200 pmol each of synthetic oligonucleotide E shown in Sequence No. 15 of the Sequence Table and synthetic

oligonucleotide F shown in Sequence No. 16 of the Sequence Table, which were designed based on the ribosomal protein L7/L12 DNA sequence of *Neisseria gonorrhoeae* acquired from Internet information

5 (Oklahoma University, *N. Gonorrhoeae* Genome Project, disclosed genome DNA data) because of the similarity with ribosomal protein L7/L12 DNA sequence of other bacteria, were added to the enzyme to bring the final volume to 50 μ l.

10 This mixture was cycled 5 times with a TaKaRa PCR Thermal Cyclor 480 for 1 minute at 95 °C, 2 minutes at 50 °C, and 3 minutes at 72 °C and was then cycled 25 times for 1 minute at 95 °C, 2 minutes at 60 °C, and 3 minutes at 72 °C. Electrophoresis was performed
15 in 1.5 % agarose gel using some of this PCR product. This product was then stained with ethidium bromide (Nihon Gene Co., ltd.) and observed under ultraviolet rays to confirm amplification of approximately 400 bp cDNA. After digestion treatment with restriction
20 endonucleases BamHI and XhoI, electrophoresis was performed in 1.5 % agarose gel and staining with ethidium bromide was carried out. An approximately 370 bp band was cut out from the gel. This band was purified with Suprec01 (Takara Co., Ltd.) and then
25 inserted into pGEX-6P-1 (Pharmacia), which is a commercial vector. This same vector can function as an expression vector for the desired molecule, which

can express fused protein with GST protein, by
insertion of the desired gene fragment into the
appropriate restriction endonuclease site. Actually,
vector pGEX-6P-1 and the previous DNA were mixed
5 together at a molar ratio of 1:5 and DNA was inserted
into the vector with T4 DNA ligase (Invitrogen Co.).
Vector pGEX-6P-1 into which DNA had been inserted was
genetically introduced to *Escherichia coli* One-Shot
Competent Cells (Invitrogen Co., Ltd.) and then
10 inoculated in a plate of L-Broth (Takara Co., ltd.)
semi-solid culture plate containing 50 µg/ml
ampicillin (Sigma). The plate was then set aside at
37 °C for 12 hours and the colonies that grew were
selected at random and inoculated into 2 ml L-Broth
15 liquid culture medium containing the same
concentration of ampicillin. Shake cultivation was
performed at 37 °C for 8 hours and the bacteria was
recovered and the plasmid was separated using Wizard
Miniprep in accordance with the attached literature.
20 The plasmid was cleaved with restriction endonuclease
BamHI/XhoI. Insertion of said PCR product was
confirmed by cutting out approximately 370 bp DNA.
The base sequence of the DNA that had been inserted
was determined using said clone.

25 Determination of the base sequence of the
inserted DNA fragment was performed using the
Fluorescence Sequencer of Applied Biosystems. The

sequence sample was prepared using PRISM, Ready Reaction Dye Terminator Cycle Sequencing Kit (Applied Biosystems). First, 9.5 μ l reaction stock solution, 4.0 μ l T7 promoter primer at 0.8 pmol/ μ l (Gibco BRL) and 6.5 μ l template DNA for sequencing at 0.16 μ g/ μ l were added to a microtube with a capacity of 0.5 ml and mixed. After layering with 100 μ l mineral oil, PCR amplification was performed for 25 cycles, where one cycle consisted of 30 seconds at 96 °C, 15 seconds at 55 °C, and 4 minutes at 60 °C. The product was then kept at 4 °C for 5 minutes. After the reaction was completed, 80 μ l sterilized pure water was added and stirred. The product was centrifuged and the aqueous layer was extracted 3 times with phenol-chloroform. Ten microliters 3 M sodium acetate (pH 5.2) and 300 μ l ethanol were added to 100 μ l the aqueous layer and stirred. The product was then centrifuged for 15 minutes at room temperature and 14000 rpm and the sediment was recovered. Once the sediment was washed with 75 % ethanol, it was dried under a vacuum for 2 minutes to obtain the sequencing sample. The sequencing sample was dissolved in formamide containing 4 μ l 10 mM EDTA and denatured for 2 minutes at 90 °C. This was then cooled in ice and submitted to sequencing.

One of the 4 clones obtained had homology of the sequence with the probe used for PCR. In addition,

DNA sequences extremely similar to the gene sequence of ribosomal protein L7/L12 gene of the other microorganisms, for example, *Haemophilus influenzae*, were discovered. The entire base sequence and the corresponding amino acid sequence of the structural gene moiety are as shown in Sequence ID No. 21 and No. 22 of the Sequence Table. This gene fragment clearly codes for *Neisseria gonorrhoeae* ribosomal protein L7/L12.

50 ml *Escherichia coli* into which expression vector had been inserted was cultivated overnight in a two-fold concentration YT medium at 37 °C. Then, 450ml of the two-fold concentration YT medium was heated at 37 °C for 1 hour. 50 ml of the *Escherichia coli* solution that had been cultivated overnight was introduced to 450ml of the aforementioned medium. After cultivation for one hour at 37 °C, 100 µl 500 mM IPTG was introduced and cultivated for 4 hours at 25 °C. The product was then recovered and introduced 250 ml each to a centrifugation tube and centrifuged for 20 minutes at 5000 rpm. The supernatant was discarded and dissolved in 25 ml each 50 mM Tris-HCl at a pH of 7.4 and Lysis buffer containing 25 % sucrose.

Furthermore, 1.25 ml 10% NP-40 and 125 µl 1 M MgCl₂ were added and transferred to a plastic tube. Sonication was performed 1 minute x 5 times while ice

cold. The product was centrifuged for 15 minutes at 12,000 rpm and the supernatant was recovered.

Next, the aforementioned supernatant was adsorbed on a glutathione sepharose column (manufactured by Pharmacia) conditioned with PBS. Then, the column was washed with PBS three times the bed volume. Elution was performed with 50 mM Tris-HCl at a pH of 8.0 containing 10 mM glutathione. The protein content in the fraction was determined by the pigment bonding method (Bradford method; BioRad Co.) and the main fraction was acquired. The main fraction was dialyzed three times against 3L PBS.

1 ml of a cleavage buffer containing 500 mM Tris-HCl (pH 7.0), 1.5 M NaCl, 10 mM EDTA, and 10 mM DTT was added to 10 ml of 1 mg/ml solution of the resulting GST fusion ribosomal protein L7/L12. 100 μ l of 2 u/ μ l PreScission Protease (manufactured by Pharmacia company) was further added and reacted at 4 °C to separate the GST moiety from ribosomal protein L7/L12.

The reaction solution was passed through a glutathione sepharose column which had been conditioned with PBS. The solution coming out from the column was recovered. One bed volume of PBS was passed through and also recovered. Purity of the purified ribosomal protein L7/L12 that was obtained was confirmed by electrophoresis to be approximately

90 %, showing that a purity satisfactory for an immunogen could be guaranteed.

First, 100 µg protein antigen of ribosomal protein L7/L12 of *Neisseria gonorrhoeae* was dissolved in 200 µl PBS and then 200 µl Freund's complete adjuvant was added and mixed and emulsification was performed. 200 µl was intraperitoneally injected to immunize mice. Then, the same emulsion antigen was intraperitoneally injected after 2 weeks, after 4 weeks, and after 6 weeks. A two-fold concentration antigen emulsion was further injected intraperitoneally after 10 weeks and after 14 weeks. The spleen was excised 3 days after the final immunization and submitted to cell fusion.

After thoroughly mixing 2×10^7 myeloma cells per 10^8 spleen cells from mice, which had been recovered aseptically, in a glass tube, the mixture was centrifuged for 5 minutes at 1500 rpm and the supernatant was discarded. The cells were thoroughly mixed.

The myeloma cells used for cell fusion were obtained by cultivation of cell strain NS-1 with an RPMI 1640 culture medium containing 10 % FCS, cultivating this product beginning 2 weeks before cell fusion using an RPMI 1640 medium containing 0.13 mM azaguanine, 0.5 µg/ml MC-210, and 10 % FCS for 1 weeks, and then further cultivating the cell strain

for 1 week with an RPMI 1640 medium containing 10 % FCS. 30 ml of RPMI 1640 culture medium 50 ml that had been kept at 37°C was added to the mixed cell sample and centrifuged at 1,500 rpm. After removal of the supernatant, 1 ml 50 % polyethylene glycol that had been kept at 37 °C was added and stirred for 2 minute. 10 ml RPMI 1640 medium kept at 37 °C was added and the solution was vigorously mixed for approximately 5 minutes as it was suctioned and evacuated from a sterile pipette.

After centrifugation for 5 minutes at 1,000 rpm and removal of the supernatant, 30 ml HAT culture medium were added to bring the cell concentration to 5×10^6 cells/ml. This mixture was stirred till uniform and then poured, 0.1 ml at a time, into a 96-well culture plate and cultivated at 37 °C under 7 % CO₂. HAT medium was added, 0.1 ml at a time, on Day 1 and at Week 1 and Week 2.

Then the cells that had produced the desired antibody were assessed by ELISA. Solutions of ribosomal protein L7/L12 of *Neisseria gonorrhoeae* dissolved in PBS containing 0.05 % sodium azide were diluted to 10 µg/ml and were poured, 100 µl at a time, into separate 96-well plates and adsorbed overnight at 4 °C. After removal of the supernatant, 200 µl 1 % bovine serum albumin solution (in PBS) were added and reacted and blocked for 1 hour at room temperature.

The supernatant was removed and the product was washed with a washing solution (0.02 % Tween 20, PBS). 100 μ l of a culture solution of fusion cells was added and reacted for two hours at room temperature.

5 The supernatant was removed and the product was further washed with a washing solution. Then, 100 μ l of a peroxidase-labeled goat anti-mouse antibody solution at 50 ng/ml was added and reacted for one hour at room temperature. The supernatant was removed and the product was washed with a washing solution. TMB (KPL) solution was added, 100 μ l at a time, and reacted for 20 minutes at room temperature. After coloration, 100 μ l 1 N sulfuric acid were added to stop the reaction. Absorbance at 450 nm was
10 determined.
15

As a result, positive wells that reacted with ribosomal protein L7/L12 were detected, confirming presence of the antibody to ribosomal protein L7/L12.

Therefore, the cells in the positive wells were recovered and cultivated with HAT medium in a 24-well plastic plate. The fused medium that had been
20 cultivated was diluted with an HT medium to a cell count of approximately 20 cells/ml. Then 50 μ l the medium was mixed with 10^6 six-week-old mouse thyroid
25 cells suspended in the HT medium in a 96-well culture plate. The cells were cultivated for 2 weeks at 37 °C under the conditions of 7 % CO₂. The antibody titer in

the culture supernatant was determined by the
aforementioned ELISA method and, the cells those
showed a positive reaction with ribosomal protein
L7/L12 were recovered. Furthermore, the same dilution
5 and cloning procedure was repeated to obtain a total
of 4 clones of hybridoma AMGC-5 to AMGC-8.

Monoclonal antibody was produced and recovered
according to standard methods, using the positive
hybridoma cells obtained as previously described.

10 Specifically, cells subcultured in RPMI 1640
medium (containing 10 % FCS) was diluted in serum-
free medium to about 2×10^5 cells/ml, 3.3×10^5
cells/ml, and 5×10^5 cells/ml in 25 cm² culture flask,
and the total was made 5 ml. After cells were grown
15 for 3 to 5 days in 7 % CO₂ at 37 °C, flask that
contains the least number of original cells was
selected among flasks in which cells were grown. The
same procedure was repeated until the cells diluted
to 2×10^5 cells/ml grow to 2×10^6 cells/ml in 3 to 4
20 days, thereby adapting the cells with serum-free
medium. Next, cloning was performed in a 96-well
plate for bacterial culture to select cells
exhibiting fastest growth and a highest antibody
titer. The selected cells were grown in a 24-well
25 plate and diluted with a serum-free medium in a 25
cm² culture flask to a concentration of about 2×10^5
cells/ml and the total volume was made to 10 ml.

After incubation for 3 to 4 days in 7 % CO₂ at 37 °C to a concentration of 1 x 10⁶ cells/ml, the culture broth 100 ml, 1 x 10⁶ cells/ml was transferred to a bottle for mass cultivation which were grown in the same manner in a 75 cm² flask. 100 ml of a serum-free medium was added to the mixture and was incubated at 37 °C for two days while stirring. 200 ml of the serum-free medium was added again and the mixture was incubated for a further two days. The culture broth was divided into four aliquots, one volume serum-free medium was added to each portion, followed by incubation for two days. After further addition of 400 ml of the serum-free medium, the culture broth was incubated for 6 days. The culture broth was collected and centrifuged at 10,000 rpm for 15 minutes to obtain culture supernatant containing the target antibody. After adding sodium azide to final concentration 0.1 %, the culture supernatant was stored at 4 °C. 100 ml of the supernatant containing the antibody was diluted 5-fold with PBS and adsorbed in a protein G column (5 ml bed, Pharmacia) and washed with 3-bed volume of PBS. Then eluted with citrate buffer at pH 3, the antibody fraction was recovered and monoclonal antibody produced by each hybridoma was obtained.

The monoclonal antibodies originating from the four hybridoma clones were evaluated according to the

OIA method described in International Patent
Application Japanese Laid-open No. 509565/1995.

Specifically, the OIA method comprises preparing
a reactive substrate by reacting an antibody for
5 capture on a silicon wafer having a thin film layer
of silicon nitride, causing this substrate to react
with an antigen which is an extract of microorganisms
for a prescribed period of time, causing the captured
antigen to react with an antibody (an amplification
10 reagent) which is an enzyme-labeled antibody, and
finally adding a substrate solution to produce a
thin-film precipitate. The antigen-antibody reaction
can be judged visually by a degree of light
interference color produced in the precipitate.

15 The monoclonal antibody preparation was used as a
capture antibody to be immobilized on a silicon wafer
having a silicon nitride thin film layer in the OIA
method. Moreover, peroxidase-labeled AMGC-1
monoclonal antibody which can non-specifically react
20 with ribosomal proteins L7/L12 protein of a variety
of microorganisms described in Reference Example was
used as the detect antibody. That is, enzyme
labeling was performed in accordance with the method
in "Analytical Biochemistry" 132 (1983), 68-73 with
25 the reagent S-acetylthioacetic acid N-
hydroxysuccinimide for binding using horseradish
peroxidase (Sigma Grade VI).

In the OIA reaction, monoclonal antibody in a PBS containing 0.05 % sodium azide was diluted with 0.1 M HEPES (pH 8.0) to a concentration of 10 µg/ml and added onto a silicone wafer which has a thin film layer of silicon nitride, 50 µl at a time, to react for 30 minutes at room temperature, followed by washing with distilled water and use. 15 µl of antigen solution, which had been obtained by adding 0.5 % Triton X-100 to culture suspension of various species of microorganisms and then extracting the suspension for 5 minutes at room temperature, was added to the specimen obtained in the above-described procedure and reacted for 10 minutes at room temperature. Then, 15 µl of 20 µg/ml peroxidase-labeled AMGC1 was added and reacted for 10 minutes. After washing with distilled water, a substrate solution (KPL Co.) was added, 15 µl at a time, and reacted for 5 minutes at room temperature. The product was washed with distilled water to judge the concentration of detection signals as an intensity of light interference by naked eyes.

As a result, as shown in Table 4 it is clear that when monoclonal antibody derived from hybridoma AMGC-8 was used as the capture antibody, all strains of *Neisseria* genus tested were detected at a sensitivity of 10^8 cells/ml, while reactivity of other microorganisms could not be detected. Thus, the

antibody with specific reactivity to *Neisseria* genus was confirmed to have been obtained by using the monoclonal antibody to ribosomal protein L7/L12.

5 Table 4

		Results of Detection (10 ⁸ cells/ml)
<i>Neisseria gonorrhoeae</i>	IID821	+
<i>Neisseria lactamica</i>	ATCC23970	+
<i>Neisseria meningitidis</i>	ATCC13090	+
<i>Escherichia coli</i>	ATCC25922	—
<i>Enterococcus faecalis</i>	ATCC19433	—
<i>Haemophilus influenzae</i>	ATCC10211	—
<i>Klebsiella pneumoniae</i>	ATCC13883	—
<i>Pseudomonas aeruginosa</i>	ATCC27853	—
<i>Group B streptococcus</i>	ATCC12386	—
<i>Staphylococcus aureus</i>	ATCC25923	—
<i>Streptococcus pneumoniae</i>	ATCC27336	—
<i>Streptococcus pyogenes</i>	ATCC19615	—

(+: Positive; -: Negative)

10 Example 10

Acquisition of a polyclonal antibody which specifically reacts with ribosomal protein L7/L12 of *Haemophilus influenzae* using a ribosomal protein L7/L12 protein-immobilized affinity column

15 A centrifugal supernatant of *Haemophilus influenzae* cell extract, which had been treated with 0.5% Triton X-100 was used as an antigen. About 1.2 ml of a physiological saline solution containing 100

µg of antigen was emulsified with the addition of 1.5 ml of Freund's adjuvant. The emulsion was subcutaneously injected into four SPF Japanese White Rabbits to immunize the animals. The rabbits were
5 immunized 5 to 6 times, once every two weeks, and the antibody titer was confirmed.

The antibody titer was confirmed by the ELISA method. Solutions of ribosomal protein L7/L12 of *Haemophilus influenzae* dissolved in PBS containing
10 0.05 % sodium azide diluted to 10 µg/ml were poured, 100 µl at a time, into 96-well plates and adsorbed overnight at 4 °C. After removal of the supernatant, 200 µl 1 % bovine serum albumin solution (in PBS) were added and reacted and blocked for 1 hour at room
15 temperature. The supernatant was removed and the product was washed with a washing solution (0.02 % Tween 20, PBS). 100 µl of a solution obtained by diluting normal rabbit serum and immunized rabbit antiserum was added and reacted for two hours at room
20 temperature. The supernatant was removed and the product was further washed with a washing solution. Then, 100 µl of a peroxidase-labeled goat anti-rabbit IgG antibody solution at 50 ng/ml was added and reacted for one hour at room temperature. The
25 supernatant was removed and the product was washed with a washing solution. OPD solution (Sigma Co.) was added, 100 µl at a time, and reacted for 20

minutes at room temperature. After coloration, 100 μ l of 1 N sulfuric acid was added to stop the reaction. Absorbance at 492 nm was determined.

After confirming that the antibody titer had increased, a large quantity of blood was collected. Blood was collected in a centrifugal tube made of glass from the ear artery, allowed to stand for one hour at 37 °C, then overnight at 4 °C. The mixture was centrifuged at 3000 rpm for 5 minutes and the supernatant was recovered. The resulting 4 lots of anti-serum were stored at 4 °C.

Next, an affinity column with immobilized ribosomal protein L7/L12 of *Haemophilus influenzae* and *Neisseria gonorrhoeae* was prepared. HiTrap NHS-activated column (1 ml, manufactured by Pharmacia) was used. Immediately after replacing the column with 6 ml of 1 mM HCl, 1 ml of a PBS solution of ribosomal protein L7/L12 adjusted to 1 mg/ml was charged. The column was allowed to stand for 15 minutes at 25 °C. This procedure was repeated 5 times, thereby feeding the total 5 ml of the PBS solution of ribosomal protein L7/L12. Then, 6 ml of Buffer A (0.5 M ethanolamine, 0.5 M NaCl, pH 8.3), 6 ml of Buffer B (0.1 M acetic acid, 0.5 M NaCl, pH 4), and 6 ml of Buffer A were charged as blocking reagents. After allowing to stand for 15 minutes at 25 °C, 6 ml of Buffer B, 6 ml of Buffer A, and 6 ml

of Buffer B were further added. The mixture was then equilibrated with 6 ml of PBS.

Using the affinity column with immobilized ribosomal protein L7/L12 of *Haemophilus influenzae*,
5 the polyclonal antibody in the resulting anti-serum was purified using the supernatant of Triton X-100 treated bacteria of *Haemophilus influenzae* as an antigen. This antiserum was first diluted with PBS to a volume of 5 times, passed through a 0.45 μ m filter,
10 then caused to be adsorbed in the column immobilized with ribosomal protein L7/L12 of *Haemophilus influenzae* at a flow rate of 0.5 ml/min. After elution from the column with 0.1 M glycine (pH 2.1) and immediately neutralizing with 1 M Tris-HCl (pH
15 9.0), eluted fractions of the target antibody were recovered by the ELISA method, the same as anti-titer measuring method. These fractions were passed through the affinity column immobilized with the ribosomal protein L7/L12 of *Neisseria gonorrhoeae*,
20 which was equilibrated with PBS whereby the antibody that reacts with the ribosomal protein L7/L12 of *Neisseria gonorrhoeae* was adsorbed and fraction which passed through without adsorption was recovered.

The polyclonal antibody purified in this manner
25 was evaluated by the same OIA method as in Example 6.

The purified antibody was used as a capture antibody for the OIA method. Moreover, peroxidase-

labeled AMGC-1 monoclonal antibody described in Reference Example was used as the detect antibody. That is, enzyme labeling was performed in accordance with the method in "Analytical Biochemistry" 132 (1983), 68-73 with the reagent S-acetylthioacetic acid N-hydroxysuccinimide for binding using horseradish peroxidase (Sigma Grade VI).

In the OIA reaction, the purified polyclonal antibody in a PBS containing 0.05 % sodium azide was diluted with 0.1 M HEPES (pH 8.0) to a concentration of 10 µg/ml and added onto a silicone wafer, 50 µl at a time, to react for 30 minutes at room temperature, followed by washing with distilled water and use.

15 µl of antigen solution, which had been obtained by adding 0.5 % Triton X-100 to culture suspension of various species of microorganisms and then extracting the suspension for 5 minutes at room temperature, was added to the specimen obtained in the above-described procedure and reacted for 10 minutes at room temperature. Then, 15 µl of 20 µg/ml peroxidase-labeled AMGC1 was added and reacted for 10 minutes. After washing with distilled water, a substrate solution (KPL) was added, 15 µl at a time, and reacted for 5 minutes at room temperature. The product was washed with distilled water to observe the blue color concentration by the naked eyes.

As a result, as shown in Table 5 it is clear that

when the purified polyclonal antibody of APhi2-2 was used as the capture antibody, *Haemophilus influenzae* tested at a sensitivity of 10^8 bacteria/ml was detected, while reactivity of other microorganisms could not be detected. Thus, the antibody with specific reactivity to *Haemophilus influenzae* was confirmed to have been obtained by using the polyclonal antibody purified by an affinity column immobilized with the ribosomal protein L7/L12 of *Haemophilus influenzae*.

Table 5

Results of Detection (10^8 cells/ml)		
<i>Haemophilus influenzae</i>	ATCC10211	+
<i>Escherichia coli</i>	ATCC25922	—
<i>Enterococcus faecalis</i>	ATCC19433	—
<i>Klebsiella pneumoniae</i>	ATCC13883	—
<i>Neisseria gonorrhoeae</i>	IID821	—
<i>Neisseria lactamica</i>	ATCC23970	—
<i>Neisseria meningitidis</i>	ATCC13090	—
<i>Pseudomonas aeruginosa</i>	ATCC27853	—
Group B <i>Streptococcus</i>	ATCC12386	—
<i>Staphylococcus aureus</i>	ATCC25923	—
<i>Streptococcus pneumoniae</i>	ATCC27336	—
<i>Streptococcus pyogenes</i>	ATCC19615	—

(+: Positive; -: Negative)

Reference Example 1

Acquisition of monoclonal antibody that reacts non-specifically with ribosomal protein L7/L12 of various bacteria

5 So called sandwich assay in which an antigen is sandwiched between a capture antibody and a labeled antibody for detection is useful to detect microorganisms by optical immunoassay and the ELISA method because of its high detection sensitivity. In this instance, not only an antibody which
10 specifically reacts with antigens originating from the subject microorganism, but also another antibody which recognizes an antigen epitope different from that of the specific antibody are required.

Antibodies which react non-specifically with
15 ribosomal protein L7/L12 originating various microorganisms are very useful as an antibody which can constitute a sandwich assay with an antibody which reacts specifically with ribosomal protein L7/L12.

20 Fortunately, ribosomal proteins L7/L12 protein of a variety of microorganisms have a region wherein the amino acid sequence is homologous. Here, the inventors have been successful in acquiring a monoclonal antibody which exhibits a cross reaction
25 with ribosomal proteins L7/L12 protein of various species of microorganisms from *Neisseria gonorrhoeae*. It has been discovered that an antibody to anti-

ribosomal protein L7/L12 having no specificity, which was acquired from one species of microorganisms can be used for sandwich assay of several kinds of microorganisms.

5 Cloning of ribosomal protein L7/L12 genes from *Neisseria gonorrhoeae*, mass expression in *Escherichia coli* and purification of the same protein, and preparation of monoclonal antibody to the same protein were performed.

10 After inoculating an appropriate amount of *Neisseria gonorrhoeae* strain IID821 (obtained from Tokyo University School of Medicine Laboratories) in a chocolate agar culture medium, the strain was cultivated for 24 hours in a CO₂ incubator under
15 conditions of 37 °C and 0.5 <5.0> % CO₂. The colonies that grew were suspended in a TE buffer to a final concentration of approximately 5 x 10⁹ CFU/ml. Approximately 1.5 ml of this suspension was transferred to a microcentrifugation tube and
20 centrifuged for 2 minutes at 10,000 rpm. The supernatant was discarded. The sediment was resuspended in 567 µl TE buffer. Then 30 µl 10 % SDS and 3 µl 20 mg/ml Proteinase K solution were added and thoroughly mixed. The suspension was incubated
25 for another hour at 37 °C. Next, after adding 80 µl 10 % cetyl trimethyl ammonium bromide/0.7 M NaCl solution and thoroughly mixing the product, it was

incubated for 10 minutes at 65 °C. Next, 700 µl
chloroform-isoamyl alcohol solution at a volume ratio
of 24:1 was added and stirred well. The solution was
centrifuged for 5 minutes (while being kept at 4 °C)
5 at 12,000 rpm using a microcentrifugation device and
the aqueous fraction was transferred to a new
microtube. Isopropanol was added to the fraction at
0.6-times its volume and the tube was vigorously
shaken to form sediment of the DNA. The white DNA
10 sediment was scooped with a glass rod and transferred
to a different microcentrifugation tube containing 1
ml 70 % ethanol (cooled to -20 °C).

Next, the product was centrifuged for 5 minutes
at 10,000 rpm and the supernatant was gently removed.
15 Then another 1 ml 70 % ethanol was added and the
product was centrifuged for 5 more minutes. Once the
supernatant had been removed, the sediment was
dissolved in 100 µl TE buffer to obtain the DNA
solution. The concentration of the genomic DNA
20 solution was determined quantitatively in accordance
with E5, Spectrophotometric determination of the
amount of DNA or RNA, "Molecular cloning: A
laboratory manual," 1989, Eds. Sambrook, J., Fritsch,
E.F., and Maniatis, T., Cold Spring Harbor Laboratory
Press.

25
B29
1 PCR was performed using 10 ng of this genomic
DNA. Taq polymerase (Takara Co., Ltd., code R001A)

was employed for PCR. Then, 5 μ l of a buffer attached to enzyme, 4 μ l of dNTP mixture attached to enzyme, and 200 pmol of each of synthetic oligonucleotide E shown in Sequence No. 15 of the Sequence Table and synthetic oligonucleotide F shown in Sequence No. 16 of the Sequence Table, which were designed based on the ribosomal protein L7/L12 DNA sequence of *Neisseria gonorrhoeae* acquired from Internet information (Oklahoma University, N. Gonorrhoeae Genome Project, disclosed genomic DNA data) because of the similarity with ribosomal protein L7/L12 DNA sequence of other bacteria, were added to the enzyme to bring the final volume to 50 μ l.

This mixture was cycled 5 times with a TaKaRa PCR Thermal Cycler 480 for 1 minute at 95 °C, 2 minutes at 50 °C, and 3 minutes at 72 °C and was then cycled 25 times for 1 minute at 95 °C, 2 minutes at 60 °C, and 3 minutes at 72 °C. Electrophoresis was performed in 1.5 % agarose gel using some of this PCR product. This product was then stained with ethidium bromide (Nihon Gene Co., ltd.) and observed under ultraviolet rays to confirm amplification of approximately 400 bp cDNA. After digestion treatment with restriction endonucleases BamHI and XhoI, electrophoresis was performed in 1.5 % agarose gel and staining with ethidium bromide was carried out. An approximately

370 bp band was cut out from the gel. This band was purified with Suprec01 (Takara Co., Ltd.) and then inserted into pGEX-4T-1 (Pharmacia), which is a commercial vector. Actually, vector pGEX-4T-1 and the previous DNA were mixed together at a molar ratio of 1:3 and DNA was inserted into the vector with T4 DNA ligase (Invitrogen Co.). Vector pGEX-4T-1 into which DNA had been inserted was genetically introduced to *Escherichia coli* One-Shot Competent Cells (Invitrogen Co., Ltd.) and then inoculated in a plate of L-Broth (Takara Co., Ltd.) semi-solid culture plate containing 50 µg/ml ampicillin (Sigma). The plate was then set aside at 37 °C for 12 hours and the colonies that grew were selected at random and inoculated into 2 ml L-Broth liquid culture medium containing the same concentration of ampicillin. Shake cultivation was performed at 37 °C for 8 hours and the bacteria was recovered and the plasmid was separated using Wizard Miniprep in accordance with the attached literature. The plasmid was cleaved with restriction endonuclease BamHI/XhoI. Insertion of said PCR product was confirmed by cutting out approximately 370 bp DNA. The base sequence of the DNA that had been inserted was determined using said clone.

135
B20
1 Determination of the base sequence of the inserted DNA fragment was performed using the

Fluorescence Sequencer of Applied Biosystems. The sequence sample was prepared using PRISM, Ready Reaction Dye Terminator Cycle Sequencing Kit (Applied Biosystems). First, 9.5 μ l reaction stock solution, 4.0 μ l T7 promoter primer at 0.8 pmol/ μ l (Gibco BRL) and 6.5 μ l template DNA for sequencing at 0.16 μ g/ μ l were added to a microtube with a capacity of 0.5 ml and mixed. After superposition with 100 μ l mineral oil, PCR amplification was performed for 25 cycles, where one cycle consisted of 30 seconds at 96 °C, 15 seconds at 55 °C, and 5 minutes at 60 °C. The product was then kept at 4 °C for 4 minutes. After the reaction was completed, 80 μ l sterilized pure water was added and stirred. The product was centrifuged and the aqueous layer was extracted 3 times with phenol-chloroform. Ten microliters 3 M sodium acetate (pH 5.2) and 300 μ l ethanol were added to 100 μ l aqueous layer and stirred. The product was then centrifuged for 15 minutes at room temperature and 14,000 rpm and the sediment was recovered. Once the sediment was washed with 75 % ethanol, it was dried under a vacuum for 2 minutes to obtain the sequencing sample. The sequencing sample was dissolved in formamide containing 4 μ l 10 mM EDTA and denatured for 2 minutes at 90 °C. This was then cooled in ice and submitted to sequencing. One of the 5 clones obtained had homology of the sequence with the probe

used for PCR. In addition, DNA sequences extremely similar to the gene sequence of ribosomal protein L7/L12 gene of the other microorganisms, for example, *Haemophilus influenzae*, were discovered. The entire base sequence and the corresponding amino acid sequence of the structural gene moiety are as shown in Sequence No. 21 and No. 22 of the Sequence Table. This gene fragment clearly codes for *Neisseria gonorrhoeae* ribosomal protein L7/L12.

Neisseria gonorrhoeae GST fused ribosome protein L7/L12 prepared by the same method as in Example 2 was obtained using the *Neisseria gonorrhoeae* GST fusion ribosomal protein L7/L12 expression vector constructed in this way. Furthermore, hybridoma strain AMGCl, which produces monoclonal antibody to ribosomal protein L7/L12 of *Neisseria gonorrhoeae*, was obtained in accordance with the method similar to the method of Example 3. Monoclonal antibody was produced and recovered in accordance with standard methods using the positive hybridoma cells of AMGCl strain obtained as previously described.

Specifically, cells subcultured in RPMI 1640 medium (containing 10 % FCS) was diluted with a serum-free medium to about 2×10^5 cells/ml, 3.3×10^5 cells/ml, and 5×10^5 cells/ml in 25 cm² culture flasks, and the total was made 5 ml. After cell were grown for 3 to 5 days in 7 % CO₂ at 37 °C, a flask

which contains the least number of original cells was selected among flasks in which cells were grown. The same procedure was repeated until the cells diluted to 2×10^5 cells/ml grow to 2×10^6 cells/ml in 3 to 4 days, thereby adapting the cells with the serum-free medium. Next, cloning was performed in a 96-well plate for bacteria cultivation to select cells exhibiting fastest growth and a highest antibody titer. The selected cells were grown in a 24-well plate and diluted with a serum-free medium in a 25 cm² culture flask to a concentration of about 2×10^5 cells/ml and the total volume was made 10 ml. After incubation for 3 to 4 days in 7 % CO₂ at 37 °C to a concentration of 1×10^6 cells/ml, the culture broth 100 ml, 1×10^6 cells/ml was transferred to a bottle for mass cultivation which were grown in the same manner in a 75 cm² flask. 100 ml of a serum-free medium was added to the mixture, which was incubated at 37 °C for two days while stirring. 200 ml of the serum-free medium was added again and the mixture was incubated for a further two days. The culture broth was divided into four aliquots, the serum-free medium was added to each portion, followed by incubation for two days. After further addition of 400 ml of the serum-free medium, the culture broth was incubated for 6 days. The culture broth was collected and centrifuged at 10,000 rpm for 15 minutes to obtain a

culture supernatant including the target antibody. After the addition of sodium azide to final concentration 0.1 %, the culture supernatant was stored at 4 °C. 100 ml of the solution containing the antibody that was obtained was diluted 5-fold with PBS and adsorbed in a Protein A column (5 ml bed volume, Pharmacia) and washed with 3-bed volume of PBS. Then eluted with citrate buffer at pH 3, the antibody fraction was recovered and monoclonal antibody produced by each hybridoma was obtained. The monoclonal antibody derived from the hybridoma was used in ELISA.

To evaluate the antibody, 96-well plates sensitized with ribosomal proteins L7/L12 of various microorganisms were used as an antigen. The monoclonal antibody prepared was reacted, followed by the reaction of horseradish peroxidase-labeled anti-mouse IgG (manufactured by MBL, Code 330) as a secondary antigen, and finally the antibody was detected using an enzyme reaction coloring reagent. In the ELISA reaction, solutions of recombinant ribosomal protein L7/L12 of *Neisseria gonorrhoeae*, *Haemophilus influenzae*, and *Streptococcus pneumoniae* dissolved in PBS containing 0.05 % sodium azide diluted to 1 µg/ml were poured, 100 µl at a time, into separate 96-well plates and adsorbed overnight at 4 °C. After removal of the supernatant, 200 µl 1%

bovine serum albumin solution (in PBS) were added and reacted and blocked for 1 hour at room temperature.

The supernatant was removed and the product was

washed with a washing solution (0.02 % Tween 20,

5 PBS). Solutions of AMGC1 antibody, at concentrations of 0.1-1 µg/ml, in an amount of 100 µl each, were added and reacted for two hours at room temperature.

The supernatant was removed and the product was

further washed with a washing solution. Then, 100 µl

10 of a solution of 5 µg/ml horse radish peroxidase-labeled anti-mouse IgG (manufactured by MBL, Code 330) was added and reacted for one hour at room

temperature. The supernatant was removed and the

product was washed with a washing solution. TMB (KPL)

15 solution was added, 100 µl at a time, and reacted for 20 minutes at room temperature. After coloration,

100 µl 1 N sulfuric acid were added to stop the

reaction. Absorbance at 450 nm was determined.

As a result, as shown in Table 6 it was confirmed

20 that when the monoclonal antibody originating from

hybridoma AMGC1 was used, this antibody can react

with ribosomal proteins L7/L12 of all bacteria such

as *Neisseria gonorrhoeae*, *Haemophilus influenzae*, and

Streptococcus pneumoniae.

25

Table 6

Results of detection of AMGC1 antibody and Ribosomal Protein L7/L12 of microorganisms
--

<i>Neisseria gonorrhoeae</i>	+
<i>Haemophilus influenzae</i>	+
<i>Streptococcus pneumoniae</i>	+

(+: Positive)

The AMGC1 antibody obtained here is very useful as an antibody used for the detection of microorganisms in the so-called sandwich assay by optical immunoassay and ELISA method, in combination with an anti-ribosomal protein L7/L12 antibody which is specific to each microorganism.

Industrial Applicability

According to the present invention, not only microorganisms can be detected specifically according to species, but also microorganisms of all serotypes in the same species can be detected at a high precision, by using antibodies to intracellular molecules of the same function.

By using antibodies to ribosomal proteins L7/L12 of various microorganisms as such antibodies, *Haemophilus influenzae* and *Neisseria gonorrhoeae* can be detected precisely.

Moreover, specific antibodies used for the detection of various kinds of microorganisms can be prepared by using intracellular molecules exhibiting same functions in various microorganisms as an

antigen.

REMARKS TO DEPOSITED MICROORGANISMS

The organization in which the microorganisms have been deposited:

- 5 National Institute of Bioscience and Human
 Technology, the Agency of Industrial Science
 and Technology

Address: 1-1-3, Higashi, Yatabe-cho, Tsukuba-shi,
 Ibaraki-ken, Japan (Postal Code: 305).

- 10 Date of deposition: July 28, 1999

Number of deposition given by the deposition
organization: FERM BP-6807

INS
B22